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Attribute VB_Name = "General"
Option Explicit

'Program and DB Rev
Public Const ProgRev$ = "PP Rev 1.00"
Public Const DBRev$ = "PP M03 Template 1.04"

'Columns collection
Public PSColumns As New Collection

'Statistics Xarray
Public StatArray As New Xarray

'Registry keys
Public Const SKEY_PPPPS$ = "PamelaAProbeDesigner"
Public Const VALUE_DB_TEMPLATES$ = "DBTemplate"
Public Const VALUE_DB_DIRS$ = "DBDir"
Public Const VALUE_DB_EXT$ = "DBFileExt"
Public Const VALUE_GB_DIRS$ = "GBDir"
Public Const VALUE_GB_FILE_EXT$ = "GBFileExt"
Public Const VALUE_BLAST_DIRS$ = "BLASTDir"
directory

'Error codes
Enum PPErrors
    PPERgsFileLength = 3000
    PPERgsFormat = 3001
    PPERBadRecordset = 3002
    PPERNorecordsetCreated = 3003
    PPERCancelled = 3004
    PPERdbFormat = 3005
    PPERblastVersion = 3006
    PPERFASTAFormat = 3007
    PPERseqLaddB = 3008
    PPERdLL = 3009
End Enum

'Flags to prevent sequence and probest selectors firing.
'Currently broken -- RowcolChange events are ignored.
Public AllowdbClick As Boolean

'Algorithm parameter sets
Public CreatePars As New ccCreatePars
Public PosFilterPars As New cPosFilterPars
Public LengthFilterPars As New cLengthFilterPars
Public GCFilterPars As New cGCFilterPars
Public CGDPars As New cGDPars
Public dGDFilterPars As New cGDFilterPars
Public TMPars As New cTMPars
Public TMfilterPars As New cTMfilterPars
Public dGHPars As New cDGPars
Public dGHFilterPars As New cGDFilterPars
Public dGMfilterPars As New cGDFilterPars
Public RunPars As New cRunPars
Public RunFilterPars As New cRunFilterPars
Public Progress As New cProgress
Public EntrPars As New cEntrPars

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Public TrimPars As New cTRIMPars
Public ClampPars As New cClampPars
Public ClampFilterPars As New cClampFilterPars
Public BlastPars As New cBLASTPars
Public HomologyPars As New cHomologyPars
Public HomofilterPars As New cHomofilterPars
Public ArrayPars As New cArrayPars

' SeqRecord Structure
' This structure holds read from one sequence file/DB/website.
' It is used when reading Genbank files (in which case all fields
' are filled in), or FASTA files (in which case only the Header and
' Sequence portions are filled in).
Public Type SeqRecord
    Header As String
    Locus As String
    Accession As String
    Length As Long
    Sequence As String
    End Type
End Sub

Sub Main()
    'Initialize thermodynamic parameters.
    InitThermoPars
    'start it up!
    frmMain.Show
End Sub

Attribute VB_Name = "Blast"
Option Explicit

Private Declare Function InitPH Lib "homology.dll.dll"_
    (ByVal DB$, ByVal NumProbs$, ByVal WS, ByVal MaxProbeLength$) As Long

Private Declare Function AddProbe Lib "homology.dll.dll"_
    (ByVal Probes$, ByVal ProbeLength$) As Long

Private Declare Function CalcOneHomology Lib "homology.dll.dll" (ByVal
    SeqLength$) As Long

Private Declare Function SkipSequence Lib "homology.dll" (ByVal seqLength$)
As Long

Private Declare Function GetHomology Lib "homology.dll.dll" (ByRef Homology$) As
Long

Private Declare Function TerminatePH Lib "homology.dll.dll" () As Long

Public Const MaxBLASTHeaders$ = 100

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Public Sub Calchomology(Seqs$), HP As cHomologyPars, Homology&(), RSBlast As Recordset, PName$)
'--'
'Function
' Calculate the homology of an array of probes to a database of sequences.
' Arguments
'   Seq: The sequences.
'   HP: The homology parameters.
'   Homology: The homologies.
'   RSBlast: The table of BLAST-located homologies.
'   PName: The name of the probeSet we are processing.
'--'

On Error Goto E

Dim NumSeqs      'number of sequences we are working with
Dim MaxProbesLen 'longest probe
Dim P$, S$        'Index
Dim DBFile$       'header file for database
Dim SR As SedRecord 'one header from the header file
Dim LineText$     'one line from the header file
Dim NumHeaders$  'number of headers in the header file
Dim DoHomology As Boolean 'is homology to be done for this sequence
Dim Fmt          'bit bucket
Dim N$, NumNoCheck$ 'list of accession numbers not to check homology against
Dim NoCheck$(0)    'base name of the DB files needed for homology
Dim HomologyBaseName

'Find the longest probe.
MaxProbesLen = 0
For P = 0 To NumSeqs - 1
  If Len(Seq(P)) > MaxProbesLen Then MaxProbesLen = Len(Seq(P))
Next P

'Check for the existence of the homologizer files.
HomologyBaseName = Left(HP.DB, Len(HP.DB) - 4)
For P = 1 To NumSeqs - 1
  If Filelen(HP.Path & "\\" & HomologyBaseName & ".headers") = Filelen(HP.Path & "\\" & HomologyBaseName & ".seqs")
    'Initialize the homologizer.
    If (InitPH(HP.Path & "\\" & HomologyBaseName & ".seqs", NumSeqs, HP.Seed,
      MaxProbesLen) <> 0) Then Err.Raise pperdl, "Error in InitPH DLL Call."
    'Load the probes into the homologizer index.
    For P = 0 To NumSeqs - 1
      If (AddProbe(Seq(P), Len(Seq(P))) <> 0) Then Err.Raise pperdl, "Error in AddProbe DLL Call."
    Next P
    'Open the specified DB.
    DBFile = FreeFile
    Open HP.Path & "\\" & HomologyBaseName & ".headers" For Input As #DBFile
    'Read the number of entries expected. The progressbar for this application
    'is posted with a maximum of 100, since we don't know the number of sequences

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'In advance of opening the DB.
Line Input #DBFile, LineText
NumHeaders = CLng(LineText)

'Create an array of all accessions that should not be searched against.
NumNoCheck = 0
If IsObject(RSBlast) Then
  RSBlast.MoveLast
  RSBlast.MoveFirst
  NumNoCheck = RSBlast.RecordCount
End If
ReDim NoCheck(0 To NumNoCheck)
N = 0
Do While Not RSBlast.EOF
  If Not RSBlast("Use") Then
    NoCheck(N) = RSBlast("Accession")
    N = N + 1
  End If
  RSBlast.MoveNext
Loop
NumNoCheck = N

'Homologize all the sequences in the DB.
S = 0
Do While Not EOF(DBFile)
  S = S + 1
  If (Fix(100 * S / NumHeaders) - Progress.StopAt > 0) Then
    Progress.CheckProgress Fix(100 * S / NumHeaders)
  End If
  'Read the next DB sequence.
  SR = ReadFASTAHeader(DBFile)
  'Check the header against the bad list.
  DoHomology = True
  For N = 0 To NumNoCheck - 1
    If SR.Accession = NoCheck(N) Then DoHomology = False
  Next N
  'Do it.
  If DoHomology Then
    If (Calchomology(SR.Length) <> 0) Then Err.Raise pperdl, "Error in Calchomology DLL Call."
  Else
    If (SkipSequence(SR.Length) <> 0) Then Err.Raise pperdl, "Error in SkipSequence DLL Call."
  End If
Loop

'Retrieve the answers, and clean up.
If (GetHomology(Homology(0)) <> 0) Then Err.Raise pperdl, "Error in GetHomology DLL Call."
If (TerminatePH() <> 0) Then Err.Raise pperdl, "Error in TerminatePH DLL call."
Exit Sub
E: Debug.Print "Error in Calchomology"
Err.Raise Err.Number, Err.Description
End Sub

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Public Function BLASTPSS (PSName$, ByVal Sequence$, BP As cBLASTpars,
MatchAccessions(), _MatchHeaders(), _MatchScores(), _MatchExpect$())
'Function
' Run BLAST on the sequence of a probeset, saving headers and scores of
homologues.
'Arguments
' PSName: The probeset name, used to name temporary files.
' Sequence: The base sequence of the probeset.
' BP: The BLAST search parameters.
' MatchAccession: Matching accession numbers.
' MatchHeader: The full matching header.
' MatchScores: The match scores.
' Notes
' This is a very fragile function, since it depends on the format of the
BLAST output file (checked only for version number), and relies on every
header containing an accession number (though one will be manufactured
if none is found).
'-----
On Error GoTo E

Dim BIFilenames$, BOFilenames$
Dim BIFILE$, BOFILE#
Dim BLASTbaseName$           'The root name for the BLAST files.
Dim CmDS
Dim PROCID
Dim foo
Dim LineText$, H$, GBWhere$, matchHeader

'Check for the existence of all the BLAST files. If files are absent, an error
will occur.
BLASTbaseName = Left(BP.DB, Len(BP.DB) - 4)
foo = FileLen(BP.Path & "\blast.exe")
foo = FileLen(BP.Path & "\<BLASTbaseName & ".nin")
foo = FileLen(BP.Path & "\<BLASTbaseName & ".nhr")
foo = FileLen(BP.Path & "\<BLASTbaseName & .nsq")

'BLAST file names.
BIFilename = PSName & ".fsa"
BOfilename = PSName & ".out"

'Open the BLAST input file.
BIFILE = Freefile
Open BP.Path & "\< & BIFilename For Output As BIFILE
'Write out the header.
Print #BIFILE, "> PSName = " & PSName
'Write out the sequence.
Do While Len(Sequence) > 60
Print #BIFILE, Mid$(Sequence, 1, 60)
Sequence = Right(Sequence, Len(Sequence) - 60)
Loop

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Print #BIFILE, Sequence
'Close input file.
Close BIFILE
'Put together the command string to execute.
BOfilename = PSName & ".out"
cmd = "cmd /c cd " & BP.Path
cmd = cmd & "blastall -p blastn -i " & BIFilename & " -d " & BLASTbaseName &
" -o " & BOfilename

'Run it and wait.
PROCID = Shell(cmd, vbHide)
foo = WaitOnProgram(PROCID)

'Open the output file.
BOFILE = Freefile
Open BP.Path & "\< & BOfilename For Input As #BOFILE

'Check for correct version
Line Input #BOFILE, LineText
If Left(LineText, 12) <> "BLASTN 2.0.2" Then Err.Raise ppErrBlastVersion

'Move down to MatchHeader.
On Err.Goto ErrNoSignificantMatch
Do
Line Input #BOFILE, LineText
Loop Until Instr(LineText, "sequences producing significant alignments:") <> 0
On Error Goto E

'Strip one more line
Line Input #BOFILE, LineText

'Read match information, until a blank line.
Line Input #BOFILE, LineText
H = 0
Do
LineText = Right(LineText, Len(LineText) - 68)
MatchScores(H) = CDbl(Left(LineText, 4))
LineText = Right(LineText, Len(LineText) - 4)
If IsNumeric(LineText) Then
    MatchExpectS(H) = CDbl(LineText)
Else
    MatchExpectS(H) = 0
End If
H = H + 1
Line Input #BOFILE, LineText
Loop Until Len(LineText) = 0

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Loop Until Left(LineText, 1) = ">
'Push this line onto the header.
MatchHeader(H) = LineText
'Keep pushing on left-justified lines until we see "Length".
Do
    Line Input #BOfFile, LineText
    LineText = LTrim(LineText)
    If (Left(LineText, 6) = "Length") Then Exit Do
        MatchHeader(H) = MatchHeader(H) & " " & LineText
    Loop Until False

    'Find the accession number.
    GBWhere = Instr(MatchHeader(H), "gb=")
    If (GBWhere = 0) Then
        MatchAccession(H) = "Z00000"
    Else
        GBWhere = GBWhere + 4
        Do
            MatchAccession(H) = MatchAccession(H) & Mid(MatchHeader(H), GBWhere,
1)
            GBWhere = GBWhere + 1
        Loop Until Mid(MatchHeader(H), GBWhere, 1) = " "
    End If
Next H

'Remove the BLAST files.
Close #BOfFile
Kill BP.Path & "\\" & BOfFile.Name
Kill BP.Path & "\\" & BOfFile.Name
Exit Function

ErrNoSignificantMatch:
If No match was found, so we've skipped over the match information.
    BLASTPS = 0
    Close #BOfFile
    Kill BP.Path & "\\" & BOfFile.Name
    Kill BP.Path & "\\" & BOfFile.Name
    Exit Function

BLASTPS = 0
E: Debug.Print "Error in BLASTPS"
E.Raise Err.Number, Err.Description
End Function

Attribute VB_Name = "ColumnValidation"
Option Explicit
Public Sub ColumnValidate(RSPS As Recordset)
    'Function to check the validity of columns.
    'Arguments
    '    RSPS: The parameter recordset for the Probesets being checked.
    '    Notes
    '        1. A column is valid if the values of the parameters used
    '           to calculate it are equal to the values of those parameters
    '           in the current instance of the appropriate parameter class.
    '        2. This calculation affects settings in the global state variable
    '           PScolumns. It should therefore only be called on the Probesets and
    '           Probes used to build dgProbes, i.e. dateRSPS.
    '        3. Filter columns are always valid, since they are initialized with True.
    Dim CreatePSValid As Boolean
    Dim LengthFilterValid As Boolean
    Dim PosFilterValid As Boolean
    Dim GCFFilterValid As Boolean
    Dim dGFilterValid As Boolean
    Dim TFilterValid As Boolean
    Dim dGHFilterValid As Boolean
    Dim ClampValid As Boolean
    Dim RunValid As Boolean
    Dim HomologyValid As Boolean
    Dim Col As cColumn

    'On startup, no columns exist, so bail.
    If PScolumns.Count = 0 Then Exit Sub

    'If the recordsets are empty, then nothing is valid.
    If Not IsGoodRS(RSPS) Then
        For Each Col In PSColumns
            If Col.CanBeInvalid Then Col.IsValid = False
        Next Col
        Exit Sub
    End If

    'Set everything valid.
    CreatePSValid = True
    LengthFilterValid = True
    PosFilterValid = True
    GCFFilterValid = True
    dGFilterValid = True
    TFilterValid = True
    dGHFilterValid = True
    ClampValid = True
    RunValid = True
    HomologyValid = True

    'Try to invalidate.
    RSPS.MoveFirst
    Do While Not RSPS.EOF
        CreatePSValid = CreatePSValid And CreatePSpars.Validate(RSPS)
        LengthFilterValid = LengthFilterpars.Validate(RSPS)
        PosFilterValid = PosFilterpars.Validate(RSPS)
        GCFFilterValid = GCFFilterpars.Validate(RSPS)
        dGFilterValid = dGFilterpars.Validate(RSPS)
        TFilterValid = TMFilterpars.Validate(RSPS)
        dGHFilterValid = dGHFilterpars.Validate(RSPS)
        dGDFilterValid = dGDFilterpars.Validate(RSPS)
        HomologyValid = HomologyFilterpars.Validate(RSPS)
    Loop
End Sub

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dGMValid = dGMFilterars.Validate(RSPS) And
dGPFilterars.Validate(RSPS) And
RunValid = RunFilterars.Validate(RSPS) And
RunFilterPars.Validate(RSPS) And
ClampValid = ClampPars.Validate(RSPS) And
ClampFilterPars.Validate(RSPS)
HomologyValid = HomologyPars.Validate(RSPS) And
HomologyPars.Validate(RSPS)
RSPS.MoveNext
Loop

'Update columns.
PSColumns("Sequence").IsValid = CreatePSValid
PSColumns("Length").IsValid = CreatePSValid And LengthFilterisValid
PSColumns("Length Filter").IsValid = PSColumns("Length").IsValid
PSColumns("Position").IsValid = PSColumns("Length") .IsValid
PSColumns("Pos Filter").IsValid = CreatePSValid And PosFilterisValid
PSColumns("GC").IsValid = PSColumns("Position").IsValid
PSColumns("GC Filter").IsValid = CreatePSValid And GCFilterisValid
PSColumns("Duplex dg").IsValid = PSColumns("GC").IsValid
PSColumns("DDG Filter").IsValid = dgValid
PSColumns("TM").IsValid = TMValid
PSColumns("TM").IsValid = PSColumns("Duplex dg").IsValid
PSColumns("dGH").IsValid = PSColumns("TM").IsValid
PSColumns("dGH Filter").IsValid = dGHValid
PSColumns("dGM").IsValid = PSColumns("dGM").IsValid
PSColumns("Run Length").IsValid = RunValid
PSColumns("Run Filter").IsValid = PSColumns("Run Length").IsValid
PSColumns("Clamp").IsValid = ClampValid
PSColumns("Clamp Filter").IsValid = PSColumns("Clamp").IsValid
PSColumns("Homology").IsValid = HomologyValid
PSColumns("Homology Filter").IsValid = PSColumns("Homology").IsValid

' Set column heading backcolor according to validity.
For Each Col In PSColumns
    If Col.CanBeInvalid Then
        If Col.IsValid Then
            frmMain.dbgProbes.Columns(Col.Name).HeadForecolor = vbBlack
        Else
            frmMain.dbgProbes.Columns(Col.Name).HeadForecolor = vbRed
        End If
    End If
Next Col

End Sub

Public Sub ColumnsExist(RSPS As Recordset, RSProbes As Recordset)
    'Function
    'Check the existence of columns.
    'Arguments
    '    RSPS: The probeset(s) Parameter recordset.
    '    RSProbes: The probeset probes.
    'Notes
    '    1. A column exists if
    '        A. For each probeset, the appropriate existence parameter is set.
    '        This indicates that some calculation has been performed to fill in
    '        the column, at sometime in the past. However, it is possible that

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        the selection criteria have changed since the calculation, bringing
        into scope probes that have not had the calculation performed. Thus,
        the existence parameter is necessary, but not sufficient.
        B. For each probe, the appropriate value is not NULL. A lack of
        null values is necessary and sufficient; however, it is potentially
        time-consuming to check, so condition A is checked first, and
        condition B is checked only if A passes.

    2. This calculation affects settings in the global state variable
        PSColumns. It should therefore only be called on the probesets and
        Probes used to build dbgrobes, i.e. datels and datSelQuery.

    3. Filter columns always exist, since they are initialized with True.

Dim Exists As Boolean
Dim Col As ccolumn
'On startup, no columns exist, so bail.
If PSColumns.Count = 0 Then Exit Sub
'If the recordssets are empty, then nothing exists, nothing is valid.
If Not (IsGoodRS(RSPS) And IsGoodRS(RSProbes)) Then
    For Each Col In PSColumns
        If Col.CanNotExist Then Col.Exists = False
    Next Col
    Exit Sub
End If

'Probe creation calculations.
Exists = True
RSPS.MoveFirst
Do While Not RSPS.EOF
    Exists = Exists And RSPS("CreatePS-Exists")
    RSPS.MoveNext
Loop
PSColumns("Accession").Exists = Exists
PSColumns("PName").Exists = Exists
PSColumns("Sequence").Exists = Exists
PSColumns("Length").Exists = Exists
PSColumns("Position").Exists = Exists
PSColumns("LongName").Exists = Exists
PSColumns("Long").Exists = Exists
PSColumns("TM-Exists")
PSColumns("GC").Exists = Exists

' TM Calculations.
Exists = True
RSPS.MoveFirst
Do While Not RSPS.EOF
    Exists = Exists And RSPS("TM-Exists")
    RSPS.MoveNext
Loop
If Exists = True Then
    RSProbes.FindFirst "TM = NULL"
    If Not RSProbes.NoMatch Then Exists = False
End If
PSColumns("TM").Exists = Exists

'dGH Calculations.
Exists = True
RSPS.MoveFirst
Do While Not RSPS.EOF

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    Exists = Exists And RSPS("dGH-Exists")
    RSPS.MoveNext
  Loop
  If Exists = True Then
    RSProbes.FindFirst "dGH" = NULL
    If Not RSProbes.Nomatch Then Exists = False
    End If
    PSColumns("dGH").Exists = Exists
    'dGM Calculations.
    Exists = True
    RSPS.MoveFirst
    Do While Not RSPS.EOF
      Exists = Exists And RSPS("dGM-Exists")
    Loop
    If Exists = True Then
      RSProbes.FindFirst "dGM" = NULL
      If Not RSProbes.Nomatch Then Exists = False
    End If
    PSColumns("dGM").Exists = Exists
    'Clamp Calculations.
    Exists = True
    RSPS.MoveFirst
    Do While Not RSPS.EOF
      Exists = Exists And RSPS("Clamp-Exists")
    Loop
    If Exists = True Then
      RSProbes.FindFirst "Clamp" = NULL
      If Not RSProbes.Nomatch Then Exists = False
    End If
    PSColumns("Clamp").Exists = Exists
    'Run Length Calculations.
    Exists = True
    RSPS.MoveFirst
    Do While Not RSPS.EOF
      Exists = Exists And RSPS("Run-Exists")
    Loop
    If Exists = True Then
      RSProbes.FindFirst "Run Length" = NULL
      If Not RSProbes.Nomatch Then Exists = False
    End If
    PSColumns("Run Length").Exists = Exists
    'Homology Calculations.
    Exists = True
    RSPS.MoveFirst
    Do While Not RSPS.EOF
      Exists = Exists And RSPS("Homology-Exists")
    Loop
    If Exists = True Then
      RSProbes.FindFirst "Homology" = NULL
      If Not RSProbes.Nomatch Then Exists = False
    End If
  End If
  RSPS.MoveNext
End If

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  End If
  PSColumns("Homology").Exists = Exists
  End Sub
  -----
  Attribute VB_Name = "CreateProbeset"
  Option Explicit
  Public Function CreatePSEqualIfs (seq$, CPSP As CCreatersPars, Probs$(), _
  Positions(), GC() )
  'Function:
  '  Create a set of probes using the EqualLength method.
  'Parameters:
  '  seq: The sequence from which the probes are created.
  '  CPSP: An instance of the parameter class for this algorithm.
  '  Probes: The array in which the probes will be returned.
  '  Positions: The array in which the starting positions will be returned.
  '  GC: The array in which GC content will be returned.
  'Returns:
  '  The number of probes created.
  'Notes:
  '  2. The calling routine is responsible for adequately dimensioning Probes,
  '     Positions, and GC.
  'Dim StartPos&           'the starting position of a probe
  'Dim ProbeNum&           'the current probe
  StartPos = 1
  ProbeNum = LBound(Probes)
  Do While StartPos <= Len(seq) - CPSP.Length + 1
    If ProbeNum - Progress.StopAt = 0 Then Progress.CheckProgress StartPos
    Probes(ProbeNum) = Mid(seq, StartPos, CPSP.Length)
    Positions(ProbeNum) = StartPos
    GC(ProbeNum) = DNA_GCContent(Probes(ProbeNum))
    StartPos = StartPos + CPSP.spacing
    ProbeNum = ProbeNum + 1
  Loop
  CreatePSEqualI = ProbeNum - 1
  End Function
  Public Function CreatePSEqualLNs (seq$, CPSP As CCreatersPars, Probs$(), _
  Positions(), GC() )
  'Function:
  '  Create a set of probes using the EqualLN method.
  'Parameters:
  '  seq: The sequence from which the probes are created.
  '  CPSP: An instance of the parameter class for this algorithm.
  '  Probes: The array in which the probes will be returned.
  '  Positions: The array in which the starting positions will be returned.
  '  GC: The array in which GC content will be returned.
  'Returns:
  '  The number of probes created.
  'Notes:
  '  2. The calling routine is responsible for adequately dimensioning Probes,

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        Positions, and GC.

        If (Mid$(Seq, 1, 1) = "g" Or Mid$(Seq, 1, 1) = "c") Then DNA_GCContent =
        DNA_GCContent + 1
    Next 1
    DNA_GCContent = DNA_GCContent / Len(Seq) * 100
End Function

Public Function DNA_Seqrtrim$(seq$, Bases$, WhichEnd$)
    'Function
    'Trim off one end or the other of the sequence.
    'If Bases < Len(Seq) Then
    '    If WhichEnd = "5" Then DNA_Seqrtrim = Right$(Seq, Len$(seq) - Bases)
    '    If WhichEnd = "3" Then DNA_Seqrtrim = Left$(Seq, Len$(seq) - Bases)
    End If
End Function

Public Function DNA_SeekKeep$(seq$, Bases$, WhichEnd$)
    'Function
    'Remove all but the specified number of bases.
    'If Bases <= Len$(seq) Then
    '    If WhichEnd = "5" Then DNA_SeekKeep = Left$(Seq, Bases)
    '    If WhichEnd = "3" Then DNA_SeekKeep = Right$(Seq, Bases)
    End If
End Function

Public Function DNA_RevComp$(ByVal Seq$)
    'Function
    'Given 5'-3' strand, returns 3'-5' complement.
    'Revision
    '28-Jul-97: From PK's routine of the same name.
    Dim 16
    Seq = LCase$(Seq)
    For 1 = Len$(Seq) To 1 Step -1
        Select Case Mid$(Seq, 1, 1)
        Case "a"
            DNA_RevComp = DNA_RevComp & "t"
        Case "c"
            DNA_RevComp = DNA_RevComp & "g"
        Case "g"
            DNA_RevComp = DNA_RevComp & "c"
        Case "t"
            DNA_RevComp = DNA_RevComp & "a"
        Case "u"
            DNA_RevComp = DNA_RevComp & "a"
        Case "n"
            DNA_RevComp = DNA_RevComp & "n"
        Case Else
            MsgBox "Unknown base found while calculating DNA_RevComp."
        End Select
        DNA_RevComp = Seq
    Next 1
    Exit Function
End Function

Attribute VB_Name = "DNA_Manipulations"
Option Explicit

Public Function DNA_GCContent$(ByVal Seq$)
    'Function
    'Compute the GC content of a sequence.
    Dim 16
    Seq = LCase$(Seq)
    DNA_GCContent = 0
    For 1 = 1 To Len(Seq)

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Public Function DNA_Comps(ByVal Seq$)
'-----'
'Function
' Given 5'-3' strand, returns complement. This might be useless!
'-----'
Dim 16
Seq = LCase(Seq)
For 1 = 1 To Len(Seq)
  Select Case Mid$(Seq, 1, 1)
    Case "a"
      DNA_Comp = DNA_Comp & "t"
    Case "c"
      DNA_Comp = DNA_Comp & "g"
    Case "g"
      DNA_Comp = DNA_Comp & "c"
    Case "t"
      DNA_Comp = DNA_Comp & "a"
    Case "u"
      DNA_Comp = DNA_Comp & "a"
    Case "n"
      DNA_Comp = DNA_Comp & "n"
    Case Else
      MsgBox "Unknown base found while calculating DNA_Comp."
      DNA_Comp = Seq
      Exit Function
  End Select
Next 1
End Function

Public Function DNA_Rev$(ByVal Seq$)
'-----'
'Function
' Given 5'-3' strand, returns 3'-5' sequence. This might be useless!
'-----'
Dim 16
Seq = LCase(Seq)
For 1 = Len(Seq) To 1 Step -1
  Select Case Mid$(Seq, 1, 1)
    Case "a"
      DNA_Rev = DNA_Rev & "a"
    Case "c"
      DNA_Rev = DNA_Rev & "c"
    Case "g"
      DNA_Rev = DNA_Rev & "g"
    Case "t"
      DNA_Rev = DNA_Rev & "t"
    Case "u"
      DNA_Rev = DNA_Rev & "u"
    Case "n"
      DNA_Rev = DNA_Rev & "n"
    Case Else
      MsgBox "Unknown base found while calculating DNA_Rev."
      DNA_Rev = Seq
      Exit Function
  End Select
Next 1
End Function

```

```

Public Sub DNA_Str2Num(ByVal Seq$, ByRef NumSeq$(1))
'-----'
'Function
' Return a numeric array representing the DNA string.
'Arguments
'   Seq: The original sequence.
'   NumSeq: An integer array representing the sequence, w/ lower bound 0.
'Notes:
' 1. NumSeq must be correctly dimensioned by the calling routine.
'     since VB won't let me return an array from a function.
'-----'
Dim 16
Seq = LCase(Seq)
For 1 = 0 To Len(Seq) - 1
  Select Case Mid$(Seq, 1 + 1, 1)
    Case "a"
      NumSeq(1) = 0
    Case "c"
      NumSeq(1) = 1
    Case "g"
      NumSeq(1) = 2
    Case "t"
      NumSeq(1) = 3
    Case "u"
      NumSeq(1) = 3
    Case "n"
      NumSeq(1) = -1
  End Select
Next 1
MsgBox "Unknown base found while calculating DNA_Str2Num."
End Sub

Public Function DNA_Num2Str$(NumSeq$(1))
'-----'
'Function:
'   Return the string representation of the DNA string
'Arguments:
'   NumSeq: An integer array representing the sequence.
'-----'
Dim 16
For 1 = 0 To UBound(NumSeq)
  Select Case NumSeq(1)
    Case "0"
      DNA_Num2Str = DNA_Num2Str & "a"
    Case "1"
      DNA_Num2Str = DNA_Num2Str & "c"
    Case "2"
      DNA_Num2Str = DNA_Num2Str & "g"
    Case "3"
      DNA_Num2Str = DNA_Num2Str & "t"
    Case "-1"
      DNA_Num2Str = DNA_Num2Str & "n"
  End Select
Next 1
MsgBox "Unknown numeric code found while calculating DNA_Num2Str."
End Sub

```

```

End Function

Attribute VB_Name = "Engine"
Option Explicit

Private sub Pscalblast(RSPS As Recordset, RBLast As Recordset)
'Function
' Perform all database gets/puts, etc, for BLAST calculation.
' Arguments
' RSPS: The parameters recordset (set parameters used in current record).
' On Error GoTo E
    Dim PSName$           'probe set name.
    Dim Sequences$        'sequence!
    Dim MatchAccessions$ (MaxBLASTHeaders)
    Dim MatchAccessions$ (MaxBLASTHeaders)
    Dim MatchHeaders$ (MaxBLASTHeaders)
    Dim MatchScores$ (MaxBLASTHeaders)
    Dim MatchExpect$ (MaxBLASTHeaders)
    Dim H#
    Dim NumMatches#
    Dim FindStr$          'headers of matches
    Dim Scores#            'scores of matches
    Dim ExpectedValue#    'expected value of matches
    Dim LoopIndex#         'loop index
    Dim NumMatchesFound# 'number of matches found

    'Get the probe set name and sequence.
    PSName = RSPS("PSName")
    Sequence = RSPS("CreatePS-Sequence")
    If Len(Sequence) = 0 Then GoTo UpdateOnly

    'Run it.
    Progress.ShowProgress "Calculating BLAST matches for ProbeSet " & PSName, 0, 1
    NumMatches = BLASTES(PSName, Sequence, BlastPars, MatchAccession, MatchHeaders,
    MatchScores, MatchExpect$)

    'Update the results.
    For H = 0 To NumMatches - 1
        'Check for this record already being present.
        .FindFirst "PSName = " & PSName & " AND Accession = " & H
        MatchAccession(H) & =
        If .NoMatch Then
            .AddNew
        Else
            .Edit
        End If
        .Fields("PSName") = PSName
        .Fields("Accession") = MatchAccession(H)
        .Fields("Header") = MatchHeaders(H)
        .Fields("Score") = MatchScores(H)
        .Fields("Expect") = MatchExpect(H)
        If MatchScores(H) > BlastPars.Scutoff Or MatchExpect(H) <
        BlastPars.Ecutoff Then
            .Fields("Used") = False
        Else
            .Fields("Used") = True
        End If
        .Update
    Next H
End With

```

```

UpdateOnly:
    'Store parameters in the database.
    BlastPars.StoreDB RSPS

Exit Sub
E: Debug.Print "Error in Pscalblast"
Err.Raise Err.Number, , Err.Description
End Sub

Private Sub SeqCalcCreates(RSSeq As Recordset, RSProbes As Recordset, _
    RSPS As Recordset)
    'Function
    'Create a ProbeSet
    'Arguments
    ' RSSeq: The sequence recordset (positioned to current sequence).
    ' RSPS: The Probesets recordset (used to add new probes).
    ' RSProbes: The probes recordset (used to add new probes).
    ' RSPS: The Parameters recordset (used to add a new parameter record).
    'Errors Raised
    'ppErrorNoProbesCreated: Raised if no probes are generated.
    'Errors Handled
    'None.
    'Notes
    ' 1. Under normal use (e.g. when called by the GUI), this routine uses the
    '     values in CreateRSPS.
    ' 2. Only RSSeq must be a good recordset, since all the others are
    '     used only to add records.
    Dim Accession$          'current accession
    Dim Seq$                 'current sequence
    Dim SeqLength#           'current sequence length
    Dim Num$                 'number of probes for this sequence
    Dim PSName$              'new ProbeSet name
    Dim Probes$()             'temporary store for created probes
    Dim Positions$()          'temporary store for created positions
    Dim GC$()                'temporary store for GC content
    Dim NumProbes#           'number of probes created
    Dim P#                     'index variable for traversing Probes
    Dim Progress$()           'text for progress bar

    On Error GoTo E
    'Check the recordset
    If Not IsGoodRS(RSSeq) Then Exit Sub
    'Get the sequence, accession, etc.
    Accession = RSSeq("Accession")
    SeqLength = RSSeq("Sequence")
    Seq = RSSeq("Sequence")
    Num$ = RSSeq("Length")
    'New probeset name.

```

```

PSName = Accession & " PS" & CStr(NumPS)

'Create temporary store for new probes.
ReDim Probes(SeqLength)
ReDim Positions(SeqLength)
ReDim GC(SeqLength)

'Open the progress bar.
Progress.ShowProgress "Creating ProbeSet" & PSName, 0, NumProbes

'Create the probes.
Select Case CreateSPars.Method
Case "Equal"
    NumProbes = CreatePSEqual(Seq, CreateSPars, Probes, Positions, GC)
Case "EqualTM"
    NumProbes = CreatePSEqualTM(Seq, CreateSPars, Probes, Positions, GC)
End Select
If NumProbes = 0 Then Err.Raise pErrNoProbesCreated
ReDim Preserve Probes(NumProbes)
ReDim Preserve Positions(NumProbes)
ReDim Preserve GC(NumProbes)

'Add a new ProbeSet.
RSSeq.Edit
RSSeq.Fields("NumPS") = NumPS + 1
RSSeq.Update
CreateSPars.AddNewDB RSPS, Accession, PSName, SeqLength, Seq

'Create new records in the Probes table.
Progress.ShowProgress "Updating Probes in Database for ProbeSet" & PSName, 0,
NumProbes
For P = 0 To UBound(Probes) - 1
    If P = Progress.StopAt = 0 Then Progress.CheckProgress P
    RSProbes.AddNew
    RSProbes.Fields("Accession") = Accession
    RSProbes.Fields("PSName") = PSName
    RSProbes.Fields("Sequence") = Probes(P)
    RSProbes.Fields("Length") = Len(Probes(P))
    RSProbes.Fields("Position") = Positions(P)
    RSProbes.Fields("GC") = GC(P)
    RSProbes.Update
Next P

'Clean exit.
Exit Sub
E: Debug.Print "Error in SeqCalcCreatePS"
Err.Raise Err.Number, , Err.Description
End Sub

Public Sub SeqCalcEngine(Calcs)
'Function
    'SeqCalcEngine acts as the dispatcher for calculations
    'performed on sequences. Each of the known calculations
    'can be requested by specifying its name and subname.
    'SeqCalcEngine then retrieves necessary records from the
    'DB, then calls the specified calculation for each probe.
    'Arguments

```

```

    ' Calc: The calculation to perform.
    'Notes
    ' 1. The parameters used for a particular calculation are
    ' either made available in the optional parameter array
    ' 2. This routine loops over all selected Sequences.
    'Errors
    ' Errors are raised by not handled. Calling routine is responsible
    '-----'
    Dim RSSelSeqs As Recordset          ' holds all selected Sequences
    Dim RSP As Recordset                ' holds all probes.
    Dim RSAllProbes As Recordset        ' holds all probes in one ProbeSet
    Dim RSPS As Recordset               ' holds parameters for the current ProbeSet
    Dim PSName$                         ' current ProbeSet name
    Dim ErrNumber                       ' saves number of errors raised by callee's
    Dim ErrDescription                 ' save description of errors raised by
    callee's

    'Add a new ProbeSet.
    Set RSSelSeqs = frmMain.datSelSeqs.Recordset
    If Not IsGoodRS(RSSelSeqs) Then Exit Sub
    'Check that some sequences are available.
    'Map recordset pointers.
    Set RSP = frmMain.datP.Recordset
    Set RSPS = frmMain.datPS.Recordset
    'Close the grid.
    frmMain dbgProbes.Close
    'Process all selected sequences.
    RSSelSeqs.MoveFirst
    Do While Not RSSelSeqs.EOF
        'Begin a transaction block
        'XXXX -- causes an error when entering this routine for the second time, no
        'idea why.
        'BeginTrans
        Select Case Calc
        Case "CreatePS"
            'Create the new probeset.
            SeqCalcCreatePS RSName$, RSP, RSPS
        Case "Get PS Name."
            PSName = RSPS("PSName")
        End Case
    Loop
    'Access all probes in this ProbeSet.
    With frmMain.datPonePS
        .RecordSource = "SELECT * FROM Probes WHERE PSName = " & PSName &
        .Refresh
        Set RSAllProbes = .Recordset
    End With
    '-----'

```

```

End With

'Compute length and position filters.
If LengthFilterPars.Method <> "None" Then
    PSCalcLengthFilter RSAllProbes, RSS
    PSColumns("Length Filter").IsFilter = LengthFilterPars.AppImmediate
End If
If PosFilterPars.Method <> "None" Then
    PSCalcPosFilter RSAllProbes, RSS
    PSColumns("Pos Filter").IsFilter = PosFilterPars.AppImmediate
End If
If GCFilterPars.Method <> "None" Then
    PSCalcGCFilter RSAllProbes, RSS
    PSColumns("GC Filter").IsFilter = GCFilterPars.AppImmediate
End If
PSColumns("PSName").IsVisible = True
PSColumns("Sequence").IsVisible = True
PSColumns("Length").IsVisible = True
PSColumns("GC").IsVisible = True

Case "3" Trim"
    SeqCalcTrim frmMain.datSelSeqs.Recordset, "3"
Case "5" Trim"
    SeqCalcTrim frmMain.datSelSeqs.Recordset, "5"
Case "3" Keep"
    SeqCalcKeep frmMain.datSelSeqs.Recordset, "3"
Case "5" Keep"
    SeqCalcKeep frmMain.datSelSeqs.Recordset, "5"
End Select

'End the transaction.
'CommitTrans

'Move to next selected sequence.
RSseleseqs.MoveNext

Loop

'Update appearances.
frmMain.Form_ChangeSequence
frmMain.Form_ChangerPName
'Reopen the grid.
frmMain.debugProbes.Reopen
Progress.Hide

'Clean exit.
Exit Sub

'Handle errors.
E: Debug.Print "Error in seqCalcEngine."
ErrNumber = Err.Number
ErrDescription = Err.Description

```

```

'Rollback
frmMain.Form_ChangeSequence
frmMain.Form_ChangerPName
frmMain.debugProbes.Reopen
Progress.Hide
frmMain.MousePointer = vbDefault
Err.Raise ErrNumber, , Err.Description
End Sub

Private Sub SeqCalcTrim(RS As Recordset, WhichEnd$)
    'Function
    ' Trim bases from the 3' or 5' ends.
    ' If Not ISGoodRS(RS) Then Exit Sub
    RS.Edit
    If WhichEnd = "5" Then
        RS("Sequence") = DNA_SeqTrim(RS("sequence"), TrimPars.FivePTrim, "5")
    Else
        RS("Sequence") = DNA_SeqTrim(RS("sequence"), TrimPars.ThreePTrim, "3")
    End If
    RS("Length") = Len(RS("Sequence"))
    RS("Accession") = RS("Accession") & " "
    RS.Update
    End Sub

Private Sub SeqCalcKeep(RS As Recordset, WhichEnd$)
    'Function
    ' Keep bases on the 3' or 5' ends.
    ' If Not ISGoodRS(RS) Then Exit Sub
    RS.Edit
    If WhichEnd = "5" Then
        RS("Sequence") = DNA_SeqKeep(RS("sequence"), TrimPars.FivePKeep, "5")
    Else
        RS("Sequence") = DNA_SeqKeep(RS("sequence"), TrimPars.ThreePKeep, "3")
    End If
    RS("Length") = Len(RS("Sequence"))
    RS("Accession") = RS("Accession") & " "
    RS.Update
    End Sub

Private Sub SetField(RS As Recordset, Fields, Value)
    'Function
    ' Set all entries for one field in the recordset.
    RS.MoveFirst
    Do While Not RS.EOF
        RS.Edit
        RS.Fields(Field) = Value
        RS.Update
        RS.MoveNext
    Loop
    End Sub


```

```

Private Sub PutField(RS As Recordset, Fields$, Value)
'-----'
'Function
' Put all entries for one field in the recordset.
'Notes
' Since this operation can be timeconsuming, the progressbar is updated.
' So a progressbar has to be posted first!
'-----
Dim P6
P = 0
RS.MoveFirst
Do While Not RS.EOF
  If P = Progress.StopAt = 0 Then Progress.CheckProgress P
  RS.Edit
  RS.Fields(Field) = Value(P)
  RS.Update
  P = P + 1
  RS.MoveNext
Loop
End Sub

Private Sub GetField(RS As Recordset, Fields$, Values)
'-----'
'Function
' Get all entries for one field in the recordset.
'-----
Dim P6
P = 0
RS.MoveFirst
Do While Not RS.EOF
  Values(P) = RS.Fields(Field)
  RS.MoveNext
  P = P + 1
Loop
End Sub

Private Sub PScalcDGH(RProbes As Recordset)
'-----'
'Function
' Perform all database gets/puts, etc, for dGH calculation.
'Arguments
' RProbes: The probes recordset (calculate dGH for all records).
' RSPS: The parameters recordset (set parameters used in current record).
'-----
On Error Goto E
  Dim PSName$           'name of current probeset
  Dim NumProbes$         'number of probes in the recordset
  Dim Seq$()              'sequence column from database.
  Dim dGH#()              'dGH column, to be put to database.
  'Determine the number of probes.
  NumProbes = NumRecords(RProbes)
  If NumProbes = 0 Then Goto UpdateOnly
  ' start it up.
  PSName = RSProbes("PSName")
  Progress.ShowProgress "Calculating dGH for ProbSet " & PSName, 0, NumProbes
End Sub

```

```

'Create space for database fields.
ReDim Seq(0 To NumProbes - 1)
ReDim dGH(0 To NumProbes - 1)

'Get the sequences.
GetField RSProbes, "Sequence", Seq

'Calculate dGH.
Select Case dGHParams.DR
  Case "DNA"
    DNA.CalcDGH Seq, dGHParams, dGH
  Case "RNA"
    RNA.CalcDGH Seq, dGHParams, dGH
End Select

'Update the results.
Progress.ShowProgress "Updating dGH in Database for ProbSet " & PSName, 0,
NumProbes
PutField RSProbes, "dGH", dGH

UpdateOnly:
'Store parameters in the database, and update column visibility.
dGHParams.StoreDB RSPS
PSColumns("dGH").Exists = True
PSColumns("dGH").IsVisible = True

'Calculate filter
PScalcDGHFilter RSProbes, RSPS

Exit Sub
E: Debug.Print "Error in PScalcDGH"
Err.Raise Err.Number, , Err.Description
End Sub

Private Sub PScalcDGM(RSProbes As Recordset, RSPS As Recordset)
'-----'
'Function
' Perform all database gets/puts, etc, for dGM calculation.
'Arguments
' RSProbes: The probes recordset (calculate dGM for all records).
' RSPS: The parameters recordset (set parameters used in current record).
'-----
On Error Goto E
  Dim PSName$           'name of current probeset
  Dim NumProbes$         'number of probes in the recordset
  Dim Seq$()              'sequence column from database.
  Dim dGM#()              'dGM column, to be put to database.
  'Determine the number of probes.
  NumProbes = NumRecords(RSProbes)
  If NumProbes = 0 Then Goto UpdateOnly
  ' start it up.
  PSName = RSProbes("PSName")
  Progress.ShowProgress "Calculating dGM in Database for ProbSet " & PSName, 0,
NumProbes
End Sub

```

```

Progress.ShowProgress "Calculating dGM for Probeset" & PSName, 0, NumProbes
'Create space for database fields.
ReDim Seq(0 To NumProbes - 1)
ReDim dGM(0 To NumProbes - 1)

'Get the sequences.
GetField RSprobes, "Sequence", Seq

'Calculate dGM.
Select Case dGMPars.DR
Case "DNA"
    DNA_CalcdGM Seq, dGMPars, dGM
Case "RNA"
    RNA_CalcdGM Seq, dGMPars, dGM
End Select

'Update the results.
Progress.ShowProgress "Updating dGM in Database for Probeset" & PSName, 0,
NumProbes
PutField RSprobes, "dGM", dGM
UpdateOnly:

'Store parameters in the database, and update column visibility.
dGMPars.StoreDB RSPS
RSColumns("Run Length").Exists = True
RSColumns("Run Length").IsVisible = True

'Calculate the run filter.
PSCalcRunFilter RSprobes, RSPS

Exit Sub
E: Debug.Print "Error in PSCalcRun"
Err.Raise Err.Number, , Err.Description
End Sub

Private Sub PSCalcHomology(RSprobes As Recordset, RSPS As Recordset, RSBLast As
Recordset)
'Function
'Perform all database gets/puts, etc, for Homology calculation.
'Arguments
'    RSprobes: The probes recordset (calculate Homology for all records).
'    RSPS: The parameters recordset (set parameters used in current record).
On Error Goto E

Dim PSName$                                'name of current probeset
Dim NumProbes$                             'number of probes in the recordset
Dim Seq()                                 'sequences of probes
Dim Homology()                            'run column, to be put to database

'Determine the number of probes.
NumProbes = NumRecords(RSprobes)
If NumProbes = 0 Then Goto UpdateOnly

'Start it up.
'Note that the progress bar in this case will show progress over sequences
'in the database. Since we don't know the total, we'll set the maximum to

```

```

'Start it up.
PSName = RSprobes("PSName")
Progress.ShowProgress "Calculating Run for Probeset" & PSName, 0, NumProbes
'Create space for database fields.
ReDim Pos(0 To NumProbes - 1)
ReDim Run(0 To NumProbes - 1)

'Get the sequences,
GetField RSprobes, "Position", Pos

'Calculate Run.
CalcRun Pos, RunPars, Run
UpdateOnly:
'Store parameters in the database, and update column visibility.
RunPars.StoreDB RSPS
RSColumns("Run Length").Exists = True
RSColumns("Run Length").IsVisible = True

'Calculate the run filter.
PSCalcRunFilter RSprobes, RSPS

Exit Sub
E: Debug.Print "Error in PSCalcRun"
Err.Raise Err.Number, , Err.Description
End Sub

Private Sub PSCalcHomology(RSprobes As Recordset, RSPS As Recordset, RSBLast As
Recordset)
'Function
'Perform all database gets/puts, etc, for Homology calculation.
'Arguments
'    RSprobes: The probes recordset (calculate Homology for all records).
'    RSPS: The parameters recordset (set parameters used in current record).
On Error Goto E

Dim PSName$                                'name of current probeset
Dim NumProbes$                             'number of probes in the recordset
Dim Seq()                                 'sequences of probes
Dim Homology()                            'run column, to be put to database

'Determine the number of probes.
NumProbes = NumRecords(RSprobes)
If NumProbes = 0 Then Goto UpdateOnly

'Start it up.
'Note that the progress bar in this case will show progress over sequences
'in the database. Since we don't know the total, we'll set the maximum to

```

```

'100, and have the called routine do fraction calculations.
PName = RSpores("PName")
Progress.ShowProgress "Calculating Homology for Probeset" & PName, 0, 100

'Create space for database fields.
ReDim Seq(0 To NumProbes - 1)
ReDim TM(0 To NumProbes - 1)

'Get the sequences.
GetField RSpores, "Sequence", Seq

'Calculate Homology.
CatchHomology Seq, HomologyPars, Homology, RSBlast, PName

'Update the results.
Progress.ShowProgress "Updating Homology in Database for Probeset" & PName, 0,
NumProbes
PutField RSpores, "Homology", Homology

UpdateOnly:

'Score parameters in the database, and update column visibility.
HomologyPars.StoreDB RSPS
PSColumns("Homology").Exists = True
PSColumns("Homology").IsVisible = True

'Calculate the homology filter.
PSCalchomofFilter RSpores, RSPS

Exit Sub
E: Debug.Print "Error in PSCalchomology"
Err.Raise Err.Number, , Err.Description
End Sub

Private Sub PSCalchomofFilter RSpores As Recordset)
'-----
'Function
'Perform all database gets/puts, etc, for TM calculation.
'Arguments
'    .RSpores: The probes recordset (calculate TM for all records).
'    .RSPS: The parameters recordset (set parameters used in current record).
'-----
On Error Goto E

Dim PName$                                'name of current probeset
Dim NumProbes$                            'number of probes in the recordset
Dim Seqs()                                'sequence column from database.
Dim TMs()                                 'TM column, to be put to database.
Dim dGd()

'Determine the number of probes.
NumProbes = NumRecords(RSpores)
If NumProbes = 0 Then GoTo UpdateOnly

'Start it up.
PName = RSpores("PName")
Progress.ShowProgress "Calculating TM for Probeset" & PName, 0, NumProbes

'Create space for database fields.

```

```

'Radim Seq(0 To NumProbes - 1)
'ReDim TM(0 To NumProbes - 1)

'Get the sequences.
GetField RSpores, "Sequence", Seq

'Calculate TM.
Select Case TM.Pars.Duplex
Case "DNA/DNA"
    DNA_CalcLTM Seq, TMPars, TM
Case "DNA/RNA"
    DR_CalcLTM Seq, TMPars, TM
End Select

'Update the results.
Progress.ShowProgress "Updating TM in Database for Probeset" & PName, 0,
NumProbes
PutField RSpores, "TM", TM

UpdateOnly:

'Store parameters in the database, and update column visibility.
TMPars.StoreDB RSPS
PSColumns("TM").Exists = True
PSColumns("TM").IsVisible = True

'Calculate the filter.
PSCalchomofFilter RSpores, RSPS

Exit Sub
E: Debug.Print "Error in PSCalchom"
Err.Raise Err.Number, , Err.Description
End Sub

Private Sub PSCalchomofFilter RSpores As Recordset)
'-----
'Function
'Perform all database gets/puts, etc, for TM calculation.
'Arguments
'    .RSpores: The probes recordset (calculate TM for all records).
'    .RSPS: The parameters recordset (set parameters used in current record).
'-----
On Error Goto E

Dim PName$                                'name of current probeset
Dim NumProbes$                            'number of probes in the recordset
Dim Seqs()                                'sequence column from database.
Dim dGd()                                 'dGd column, to be put to database.

'Determine the number of probes.
NumProbes = NumRecords(RSpores)
If NumProbes = 0 Then GoTo UpdateOnly

'Start it up.
PName = RSpores("PName")
Progress.ShowProgress "Calculating TM (Duplex) for Probeset" & PName, 0,
NumProbes

```

```

'Create space for database fields.
ReDim Seq(0 To NumProbes - 1)
ReDim dG(0 To NumProbes - 1)

'Get the sequences.
GetField RSprobes, "Sequence", Seq

'Calculate dCD.
Select Case dGDPars.Duplex
Case "DNA"
    DNA_CalcdG Seq, dGDPars, dGD
Case "RNA"
    DR_CalcdG Seq, dGDPars, dCD
End Select

'Update the results.
Progress.ShowProgress "Updating dG (Duplex) In Database for Probeset # & PSName, 0,
NumProbes
PutField RSprobes, "Duplex dG", dG0
updateOnly:

' store parameters in the database, and update column visibility.
dGDPars.StoreDB RSPS
PSColumns("Duplex dG").Exists = True
PSColumns("Duplex dG").IsVisible = True

'Calculate the filter.
PSCalcGDFilter RSprobes, RSPS

Exit Sub
E: Debug.Print "Error in PSCalcClamp"
Err.Raise Err.Number, , Err.Description
End Sub

Private Sub PSCalcClamp(RSprobes As Recordset, RSPS As Recordset)
'Function
' Perform all database gets/puts, etc, for Clamp calculation.
' Arguments
'   RSProbes: The probes recordset (calculate Clamp for all records).
'   RSParams: The parameters recordset (set parameters used in current record).
'   RSProbes = NumRecords(RSprobes)
On Error GoTo E

Dim PSNames$ 'name of current probeset
Dim NumProbes 'number of probes in the recordset
Dim Seq(1) 'sequence column from database.
Dim Clamp() 'Clamp column, to be put to database.

'Determine the number of probes.
If NumProbes = 0 Then GoTo UpdateOnly

'Start it up.
PSName = RSProbes("PSName")
Progress.ShowProgress "Calculating Clamp for Probeset # & PSName, 0, NumProbes
UpdateOnly:

```

```

'Create space for database fields.
ReDim Seq(0 To NumProbes - 1)
ReDim Clamp(0 To NumProbes - 1)

'Get the sequences.
GetField RSprobes, "Sequence", Seq

'Calculate Clamp.
Select Case Clamp.Pars.Duplex
Case "DNA/RNA"
    DNA_CalcClamp Seq, ClampPars, Clamp
Case "DNAXXX"
    DNA_CalcClamp Seq, ClampPars, Clamp
End Select

'Update the results.
Progress.ShowProgress "Updating Clamp In Database for Probeset # & PSName, 0,
NumProbes
PutField RSprobes, "Clamp", Clamp
updateOnly:

' store parameters in the database, and update column visibility.
ClampPars.StoreDB RSPS
PSColumns("Clamp").Exists = True
PSColumns("Clamp").IsVisible = True

'Calculate the filter.
PSCalcClampFilter RSprobes, RSPS

Exit Sub
E: Debug.Print "Error in PSCalcClamp"
Err.Raise Err.Number, , Err.Description
End Sub

Public Sub PSCalcEngine(Calc$)
'Function
' Arguments
'   Calc: The calculation to perform.
'   Notes
'       1. All numeric calculations proceed by
'           - checking that the sequences column exists,
'           - checking that the current parameters set for this calculation
'             are the same as those previously used, and, if not, nulling
'             out all previous calculations.
'           - performing the calculation.
'       2. Filter calculations proceed as above, with the preliminary step
'           of checking the existence of the column to be filtered. If it isn't
'           there, the calculation is done.
'       3. The Probes grid is closed before DB updates, then opened
'           when they are all over, to reduce screen thrashing.

```

```

4. Updates to the database are placed in transaction blocks, so
   that they can be cleanly cancelled. This should also improve
   performance on network drives.
5. Calculation is always done one ProbeSet at a time, to reduce, whenever
   possible, recalculations of values. This could probably be further improved
   by only recalculating NULL values (since any non-NULL is guaranteed good by
   the above checks).
   Errors:
   Errors are handled by rolling back pending DB transactions, cleaning up the
   UI, then re-raising the same error for the caller to deal with.
-----
```

```

Dim RSSEIPS As Recordset
Dim RSAllProbes As Recordset
Dim RSQProbes As Recordset
Filters
Dim RS As Recordset
Dim RSBlast As Recordset
Dim PSNames
Dim ErrNumber
Dim ErrDescription
Dim callees'

'Error handling.
On Error GoTo E

'Check that some ProbeSets are available, with sequence information.
Set RSSEIPS = frmMain.datSEIPS.Recordset
If Not IsObject(RSSEIPS) Then Exit Sub
If Not PSColumns("Sequence").Exists Then Exit Sub
Set RSBlast = frmEditBLAST.datBLAST.Recordset

'Close the grid.
frmMain.MousePointer = vbHourglass
frmMain.dgProbes.Close

'Post the initial progress bar.
Progress_ShowProgress "Beginning Calculation: " & calc, 0, 100

'Process all selected ProbeSets.
RSelfs.MoveFirst
Do While Not RSSEIPS.EOF

   'Get PS Name.
   PSName = RSSEIPS("PSName")

   'Access all probes in this ProbeSet.
   With frmMain.datPhones
      .RecordSource = "SELECT * FROM Probes WHERE PSName = '" & PSName & "'"
      .Refresh
      Set RSAllProbes = .Recordset
   End With

   'Access probes in this ProbeSet that pass the filters.
   With frmMain.datProbePSQuery
      .RecordSource = BuildProbePSQuery(PSName)
      .Refresh
      Set RSQProbes = .Recordset
   End With

   'Position the parameter recordset for updates.
   With frmMain.dacs
      .Recordset.FindFirst "PSName = '" & PSName & "'"
      Set RSPS = .Recordset
   End With

   'Start a transaction block.
   BeginTrans

   'Choose the operation
   Select Case Calc
      Case "dGA"
         If dGDPars.Exists(RSPPS) And Not dGDPars.Validate(RSPPS) Then SetField
            RSAllProbes, "dGA", Null
         If Not dGDFilterPars.Validate(RSPPS) Then SetField RSAllProbes, "dGD"
            Filter", True
            PSCalcDGD RSQProbes, RSPPS
      Case "dGD Filter"
         If Not PSColumns("dGD").Exists Then
            If dGDPars.Exists(RSPPS) And Not dGDPars.Validate(RSPPS) Then SetField
               RSAllProbes, "dGD", Null
         If Not dGDFilterPars.Validate(RSPPS) Then SetField RSAllProbes, "dGD"
            Filter", True
            PSCalcDGD RSQProbes, RSPPS
      Case "dTM"
         If Not TMPars.Exists(RSPPS) And Not TMPars.Validate(RSPPS) Then SetField
            RSAllProbes, "TM", Null
         If Not TMFilterPars.Validate(RSPPS) Then SetField RSAllProbes, "TM"
            Filter", True
            PSCalcTM RSQProbes, RSPPS
      Case "TM Filter"
         If Not PSColumns("TM").Exists Then
            If TMPars.Exists(RSPPS) And Not TMPars.Validate(RSPPS) Then SetField
               RSAllProbes, "TM", Null
         If Not TMFilterPars.Validate(RSPPS) Then SetField RSAllProbes, "TM"
            Filter", True
            PSCalcTM RSQProbes, RSPPS
      Case "dGH"
         If dGHPars.Exists(RSPPS) And Not dGHPars.Validate(RSPPS) Then SetField
            RSAllProbes, "dGH", Null
   End If

   'Access probes in this ProbeSet that pass the filters.
   With frmMain.datProbePSQuery
      .RecordSource = "SELECT * FROM Probes WHERE PSName = '" & PSName & "'"
      .Refresh
      Set RSQProbes = .Recordset
   End With
End If

```

```

End With

'Position the parameter recordset for updates.
With frmMain.dacs
   .Recordset.FindFirst "PSName = '" & PSName & "'"
   Set RSPS = .Recordset
End With

'Start a transaction block.
BeginTrans

'Choose the operation
Select Case Calc
   Case "dGA"
      If dGDPars.Exists(RSPPS) And Not dGDPars.Validate(RSPPS) Then SetField
         RSAllProbes, "dGA", Null
      If Not dGDFilterPars.Validate(RSPPS) Then SetField RSAllProbes, "dGD"
         Filter", True
         PSCalcDGD RSQProbes, RSPPS
   Case "dGD Filter"
      If Not PSColumns("dGD").Exists Then
         If dGDPars.Exists(RSPPS) And Not dGDPars.Validate(RSPPS) Then SetField
            RSAllProbes, "dGD", Null
      If Not dGDFilterPars.Validate(RSPPS) Then SetField RSAllProbes, "dGD"
         Filter", True
         PSCalcDGD RSQProbes, RSPPS
   Case "dTM"
      If Not TMPars.Exists(RSPPS) And Not TMPars.Validate(RSPPS) Then SetField
         RSAllProbes, "TM", Null
      If Not TMFilterPars.Validate(RSPPS) Then SetField RSAllProbes, "TM"
         Filter", True
         PSCalcTM RSQProbes, RSPPS
   Case "TM Filter"
      If Not PSColumns("TM").Exists Then
         If TMPars.Exists(RSPPS) And Not TMPars.Validate(RSPPS) Then SetField
            RSAllProbes, "TM", Null
      If Not TMFilterPars.Validate(RSPPS) Then SetField RSAllProbes, "TM"
         Filter", True
         PSCalcTM RSQProbes, RSPPS
   Case "dGH"
      If dGHPars.Exists(RSPPS) And Not dGHPars.Validate(RSPPS) Then SetField
         RSAllProbes, "dGH", Null
   End If

   'Access probes in this ProbeSet that pass the filters.
   With frmMain.datProbePSQuery
      .RecordSource = "SELECT * FROM Probes WHERE PSName = '" & PSName & "'"
      .Refresh
      Set RSQProbes = .Recordset
   End With
End If

```

```

        If Not dGHFilterPars.Validate(RSPS) Then SetField RSAllProbes, "dGH
        Filter", True
        PSCalcDGH RSQProbes, RSPS

        Case "dGH Filter"
          If Not PSColumns("dGH").Exists Then
            RSAllProbes, "dGH", Null
            If dGHPars.Exists (RSPS) And Not dGHPars.Validate(RSPS) Then SetField
              RSAllProbes, "Clamp", Null
              If Not ClampFilterPars.Validate(RSPS) Then SetField RSAllProbes, "Clamp
              Filter", True
              PSCalcClamp RSQProbes, RSPS

            Case "Clamp Filter"
              If Not PSColumns("Clamp").Exists Then
                If Not ClampPars.Exist (RSPS) And Not ClampPars.Validate(RSPS) Then
                  SetField RSAllProbes, "Clamp", Null
                  If Not ClampFilterPars.Validate(RSPS) Then SetField RSAllProbes,
                    "Clamp Filter", True
                    PSCalcClamp RSQProbes, RSPS

                Else
                  If Not ClampFilterPars.Validate(RSPS) Then SetField RSAllProbes,
                    "Clamp Filter", True
                    PSCalcClamp RSQProbes, RSPS

              End If

            Case "Length Filter"
              If Not PSColumns("Length") Exists Then Exit Sub
              If Not PSColumns("Length") Exists Then
                If Not LengthFilterPars.Validate(RSPS) Then SetField RSAllProbes,
                  "Length Filter", True
                  PSCalcLengthFilter RSQProbes, RSPS

              Case "Pos Filter"
                If Not PSColumns("Position") .Exists Then Exit Sub
                If Not PosFilterPars.Validate(RSPS) Then SetField RSAllProbes, "Pos
                Filter", True
                PSCalcPosFilter RSQProbes, RSPS

              Case "GC Filter"
                If Not PSColumns("GC") .Exists Then Exit Sub
                If Not GCFilterPars.Validate(RSPS) Then SetField RSAllProbes, "GC
                Filter", True
                PSCalcGCFilter RSQProbes, RSPS

              Case "BLAST"
                PSCalcBlast RSPS, RSBLast

              Case "Homology"
                If HomologyPars.Exists (RSPS) And Not HomologyPars.Validate(RSPS) Then
                  SetField RSAllProbes, "Homology", Null
                  If Not HomologyPars.Validate(RSPS) Then SetField RSAllProbes,
                    "Homology Filter", True
                    PSCalcHomology RSQProbes, RSPS, RSBLast

                Case "Homology Filter"
                  If Not PSColumns("Homology") .Exists Then
                    If HomologyPars.Exists (RSPS) And Not HomologyPars.Validate(RSPS)
                      Then setField RSAllProbes, "Homology", Null
                      If Not HomologyPars.Validate(RSPS) Then SetField RSAllProbes,
                        "Homology Filter", True
                        PSCalcHomology RSQProbes, RSPS, RSBLast

                  Else
                    If Not HomologyPars.Validate(RSPS) Then SetField RSAllProbes,
                      "Homology Filter", True
                      PSCalcHomology RSQProbes, RSPS, RSBLast

              End If

            Case "Run"
              If RunPars.Exists (RSPS) And Not RunPars.Validate(RSPS) Then SetField
                RSAllProbes, "Run Length", Null
                If Not RunFilterPars.Validate(RSPS) Then SetField RSAllProbes, "Run
                Filter", True
                PSCalcRun RSQProbes, RSPS

            Case "Run Filter"
              If Not PSColumns("Run Length") .Exists Then
                If RunPars.Exists (RSPS) And Not RunPars.Validate(RSPS) Then SetField
                  RSAllProbes, "Run", True
                  If Not RunFilterPars.Validate(RSPS) Then SetField RSAllProbes, "Run
                  Filter", True
                  PSCalcRun RSQProbes, RSPS

                Else
                  If Not RunFilterPars.Validate(RSPS) Then SetField RSAllProbes, "Run
                  Filter", True
                  PSCalcRunFilter RSQProbes, RSPS

              End If

            Case "Clamp"

```

```

        End If
    End Select
    'End the transaction.
    CommitTrans

    'Move to next selected probeset.
    RSSEIPS.MoveNext

    Loop

    'Reopen the grid.
    progress.Hide
    frmMain.dbgProbes.ReOpen
    DoEvents

    'Update appearances.
    frmMain.Form_ChangePSName

    'Clean exit.
    frmMain.MousePointer = vbDefault
    Exit Sub

    'Handle errors.
    E: Debug.Print "Error in PScalcEngine."
    Err.Number = Err.Number
    Err.Description = Err.Description
    Rollback
    frmMain.Form_ChangePSName
    frmMain.dbgProbes.ReOpen
    frmMain.MousePointer = vbDefault
    Err.Raise Err.Number, , Err.Description
    End Sub

    Private Sub PScalcTMFilter(RSProbes As Recordset, RSPS As Recordset)
        'Function
        'Perform all database gets/puts, etc, for TM filter calculation.
        'RSProbes: The probes recordset (calculate TM Filter for all records).
        'RSPS: The parameter recordset (set parameters used in current record).
        On Error GoTo E
        Dim PSName$           'name of current probeset
        Dim NumProbes          'number of probes in the recordset
        Dim dG#()              'dG column from database
        Dim dGDFilter() As Boolean   'dG Filter column to database
        'Determine the number of probes.
        NumProbes = NumRecords(RSProbes)
        If NumProbes = 0 Then Goto UpdateOnly

        Dim PSName$           'name of current probeset
        Dim NumProbes          'number of probes in the recordset
        Dim TM#()              'TM column from database
        Dim TMFilter() As Boolean   'TM Filter column to database
        'Determine the number of probes.
        NumProbes = NumRecords(RSProbes)
        If NumProbes = 0 Then Goto UpdateOnly

        'Start it up.
        PSName = RSProbes("PSName")
        Progress.ShowProgress "Calculating TM Filter for Probeset " & PSName, 0,
        NumProbes

        'Create space for database fields.
        ReDim TM(0 To NumProbes - 1)
        ReDim TMFilter(0 To NumProbes - 1)

        'Get the TMs.
        GetField RSProbes, "TM", TM

        'Calculate TM Filter.
        CalcTMFilter TM, TMFilterPars, TMFilter

        'Update the results.
        Progress.ShowProgress "Updating TM Filter in Database for Probeset " & PSName,
        0, NumProbes
        PutField RSProbes, "TM Filter", TMFilter

        UpdateOnly:
        'store parameters in the database, and update column visibility.
        TMFilterPars.StoreDB RSPS
        PSColumns("TM Filter").IsFilter = TMFilter.Pars.AppImmediate

        Exit Sub
        E: Debug.Print "Error in PScalcTMFilter"
        Err.Raise Err.Number, , Err.Description
        End Sub

        Private Sub PScalcGDFilter(RSProbes As Recordset, RSPS As Recordset)
        'Function
        'Perform all database gets/puts, etc, for GGD filter calculation.
        'Arguments
        '    RSProbes: The probes recordset (calculate dGD Filter for all records).
        '    RSPS: The parameter recordset (set parameters used in current record).
        On Error GoTo E
        Dim PSName$           'name of current probeset
        Dim NumProbes          'number of probes in the recordset
        Dim dG#()              'dG column from database
        Dim dGDFilter() As Boolean   'dG Filter column to database
        'Determine the number of probes.
        NumProbes = NumRecords(RSProbes)
        If NumProbes = 0 Then Goto UpdateOnly

        'Start it up.
        PSName = RSProbes("PSName")
        Progress.ShowProgress "Calculating dG (Duplex) Filter for Probeset " & PSName,
        0, NumProbes

        'Create space for database fields.
        ReDim dG(0 To NumProbes - 1)
        ReDim dGDFilter(0 To NumProbes - 1)

```

```

'Get the dgDs.
GetField RSProbes, "Duplex dG", dgD
'Calculate dGD Filter.
CalcDGDFilter dgD, dgDFilterPars, dgDFilter

'Update the results.
Progress.ShowProgress "Updating dg (Duplex) Filter in Database for ProbeSet = &
PSName, 0, NumProbes
PutField RSProbes, "Clamp Filter", ClampFilter
UpdateOnly:

'Update the results.
Progress.ShowProgress "Updating dg (Duplex) Filter in Database for ProbeSet = &
PSName, 0, NumProbes
PutField RSProbes, "dGD Filter", dgDFilter
UpdateOnly:

'Update the results.
Progress.ShowProgress "Updating dG (Duplex) Filter in Database for ProbeSet = &
PSName, 0, NumProbes
PutField RSProbes, "dGD Filter", dgDFilter
UpdateOnly:

'Update parameters in the database, and update column visibility.
'store parameters in the database, and update column visibility.
'clampfilterpars.StoreDB RSPPS
'PColumns ("clamp filter").IsFilter = ClampFilterPars.AppImmediate
'Update Only:
'Exit Sub
'E: Debug.Print "Error in PSCalcClampFilter"
'Err.Raise Err.Number, , Err.Description
End Sub
Private Sub PSCalcDGHFilter(RSProbes As Recordset, RSPPS As Recordset)
'Function
'Perform all database gets/puts, etc, for dGH filter calculation.
'Arguments
'    RSProbes: The probes recordset (calculate dGH Filter for all records)
'    RSPPS: The parameter recordset (set parameters used in current record).
'    RSPPS: The parameter recordset (set parameters used in current record).
'On Error GoTo E
Dim PSName$           'name of current probeset
Dim NumProbes#          'number of probes in the recordset
Dim DGH()               'DGH column from database
Dim DGHFilter() As Boolean
'Determine the number of probes.
NumProbes = NumRecords(RSProbes)
If NumProbes = 0 Then GoTo UpdateOnly

'Start it up.
PSName = RSProbes("PSName")
Progress.ShowProgress "Calculating dGH Filter for ProbeSet = & PSName, 0,
NumProbes

'Create space for database fields.
ReDim dGH(0 To NumProbes - 1)
ReDim DGHFilter(0 To NumProbes - 1)

'Get the dGhs.
GetField RSProbes, "dGH", dGH
'Calculate dGH Filter.
CalcDGHFilter dGH, DGHFilterPars, DGHFilter

'Update the results.
Progress.ShowProgress "Updating dGH Filter in Database for ProbeSet = & PSName,
0, NumProbes
PutField RSProbes, "dGH Filter", dGHFilter
UpdateOnly:

'store parameters in the database, and update column visibility.
'calcclampfilter.Clamp, ClampFilterPars, ClampFilter

```

```

PSColumns("dGH Filter").IsFilter = dGHFilterPars.AppImmediate

Exit Sub
E: Debug.Print "Error in PSCalcGFilter"
Err.Raise Err.Number, , Err.Description
End Sub

Private Sub PSCalcGFilter(RSProbes As Recordset, RSPS As Recordset)
'-----'
    Function
        ' Perform all database gets/puts, etc, for GC filter calculation.
        ' Arguments
            ' RSProbes: The probes recordset (calculate GC Filter for all records).
            ' RSPS: The parameter recordset (set parameters used in current record).
        '-----'
    On Error GoTo E

        Dim PSName$           'name of current Probeset
        Dim NumProbes#         'number of probes in the recordset
        Dim GC()               'GC column from database
        Dim GCFilter() As Boolean
        '-----'

        'Determine the number of probes.
        NumProbes = NumRecords(RSProbes)
        If NumProbes = 0 Then GoTo UpdateOnly

        'Start it up.
        PSName = RSProbes("PSName")
        Progress.ShowProgress "Calculating GC Filter for Probeset " & PSName, 0,
        NumProbes

        'Create space for database fields.
        ReDim GC(0 To NumProbes - 1)
        ReDim GCFilter(0 To NumProbes - 1)

        'Get the GCs.
        GetField RSProbes, "GC", GC

        'Calculate GC Filter.
        CalcGCFilter GC, GCFilterPars, GCFilter

        'Update the results.
        Progress.ShowProgress "Updating GC Filter in Database for Probeset " & PSName,
        0, NumProbes
        PutField RSProbes, "GC Filter", GCFilter

        UpdateOnly:

        'Store parameters in the database, and update column visibility.
        GCFilters.StoredDB.RSPS
        PSColumns("GC Filter").IsFilter = GCFilterPars.AppImmediate

    Exit Sub
E: Debug.Print "Error in PSCalcGFilter"
Err.Raise Err.Number, , Err.Description
End Sub

Private Sub PSCalcGDFilter(RSProbes As Recordset, RSPS As Recordset)
'-----'

```

```

'Function
' Perform all database gets/puts, etc, for dGM filter calculation.
'Arguments
'   RSProbes: The probes recordset (calculate dGM Filter for all records).
'   RSPS: The parameter recordset (set parameters used in current record).
'-----[On Error Goto E]-----[Dim PSName$]-----[Dim NumProbes#]
'-----[Dim dGM#]-----[Dim dGMFilter() As Boolean]-----[Dim dGMColumn From Database]
'-----[dGM Filter column to database]-----[Dim NumProbes = NumRecords(RSProbes)]
'-----[If NumProbes = 0 Then GOTO UpdateOnly]-----[PSName = RSProbes("PSName")]
'-----[Progress ShowProgress "Calculating dGM Filter for ProbeSet " & PSName, 0]
'-----[NumProbes]-----[Start It Up]
'-----[Create space for database fields.]-----[Redim dGM(0 To NumProbes - 1)]
'-----[Redim dGMFilter(0 To NumProbes - 1)]-----[Get the dGMs.]
'-----[GetField RSProbes, "dGM", dGM]
'-----[Calculate dGM Filter.]-----[CALCGMFilter dGM, dGMFilterPars, dGMFilter]
'-----[Update the results.]-----[Progress ShowProgress "Updating dGM Filter in Database for ProbeSet " & PSName,
'-----[0, NumProbes]-----[PutField RSProbes, "dGM Filter", dGMFilterPars, AppImmediate
'-----[UpdateOnly:]-----[StoRe Parameters In The Database, And Update Column Visibility.
'-----[dGMFilterPars, StoReB RSPS]-----[PSColumns("dGM Filter").ISFilter = dGMFilterPars.AppImmediate
'-----[Exit Sub]-----[Private Sub PSCalcRunFilter(RSProbes As Recordset, RSPS As Recordset)
'-----[E: Debug Print "Error in PSCalcDGMFilter"]
'-----[Err.Raise Err.Number, , Err.Description]
'-----[End Sub]-----[Function
'-----[Perform all database gets/puts, etc, for Run filter calculation.
'-----[Arguments
'-----[RSProbes: The probes recordset (calculate Run Filter for all records).
'-----[RSPS: The parameter recordset (set parameters used in current record).
'-----[On Error Goto E]]-----[-----]
```

```

Dim PSName$           'name of current probeset
Dim NumProbes&        'number of probes in the recordset
Dim Run()              'Run column from database
Dim RunFilter() As Boolean 'Run Filter column to database

'Determine the number of probes.
NumProbes = NumRecords(RSProbes)
If NumProbes = 0 Then Goto UpdateOnly

'Start it up.
PSName = RSProbes("PSName")
Progress.ShowProgress "Calculating Homology Filter for Probeset " & PSName, 0,
NumProbes

'Dim Run(0 To NumProbes)          'Run column from database
Dim RunFilter(0 To NumProbes - 1) 'Run Filter column to database

'Create space for database fields.
Redim Homology(0 To NumProbes - 1)
Redim HomoFilter(0 To NumProbes - 1)

'Get the Homology.
GetField RSProbes, "Homology", Homology

'Calculate Homology Filter.
CalcHomofilter Homology, HomoFilterPars, HomoFilter

'Update the results.
Progress.ShowProgress "Updating Homology Filter in Database for Probeset " &
PSName, 0, NumProbes
PutField RSProbes, "Homology Filter", HomoFilter

UpdateOnly:

'Store parameters in the database, and update column visibility.
HomoFilterPars.StoreDB RSProbes
PSColumns("Homology Filter").IsFilter = HomoFilterPars.AppImmediate

Exit Sub
E: Debug.Print "Error in PSCalchomofilter"
Err.Raise Err.Number, , Err.Description
End Sub

Private Sub PSCalcLengthFilter(RSProbes As Recordset, RSPS As Recordset)
'Function
'Perform all database gets/puts, etc, for Length filter calculation.
'Arguments
'    RSProbes: The probes recordset (calculate length filter for all records).
'    RSPS: The parameter recordset (set parameters used in current record).
'    -----
On Error Goto E
    'name of current probeset
    Dim PSName$           'name of current probeset
    Dim NumProbes&        'number of probes in the recordset
    Dim Length&()          'Length column from database
    Dim LengthFilter() As Boolean 'Length filter column to database

    'Determine the number of probes.
    NumProbes = NumRecords(RSProbes)
    If NumProbes = 0 Then Goto UpdateOnly

    'start it up.
    PSName = RSProbes("PSName")
    Progress.ShowProgress "Calculating Length Filter for Probeset " & PSName, 0,
    NumProbes

    'Determine the number of probes.
    NumProbes = NumRecords(RSProbes)

```

```

'Create space for database fields.
ReDim Length(0 To NumProbes - 1)
ReDim LengthFilter(0 To NumProbes - 1)

'Get the Lengths.
GetField RSprobes, "Length", Length

'Calculate Length Filter.
CalcLengthFilter Length, LengthFilterPars, PosFilter
CalcLengthFilter Length, LengthFilterPars, LengthFilter

'Update the results.
Progress.ShowProgress "Updating Length Filter in Database for Probeset " & PSName,
PSName, 0, NumProbes
PutField RSprobes, "Length Filter", LengthFilter
PutField RSprobes, "Pos Filter", PosFilter
UpdateOnly:

'Update the results.
Progress.ShowProgress "Updating Length Filter in Database for Probeset " &
PSName, 0, NumProbes
PutField RSprobes, "Length Filter", LengthFilter
PutField RSprobes, "Pos Filter", PosFilter
UpdateOnly:

'Get the sequence length.
SeqLength = RSPS("CreatePS-SeqLength")

'Calculate Pos Filter.
CalcPosFilter Pos, SeqLength, PosFilterPars, PosFilter

'Update the results.
Progress.ShowProgress "Updating Pos Filter in Database for Probeset " & PSName,
0, NumProbes
PutField RSprobes, "Pos Filter", PosFilter
UpdateOnly:

'Store parameters in the database, and update column visibility.
PosFilterPars.StoreDB RSPS
PSColumns("Pos Filter").IsFilter = PosFilterPars.AppImmediate

Exit Sub
E: Debug.Print "Error in PSCalcPosFilter"
Err.Raise Err.Number, , Err.Description
End Sub

-----
Attribute VB_Name = "Filters"
Option Explicit

Public Sub CalcTMFilter(TM(), TMFPars As CTMFfilterPars, Filter() As Boolean)
    'Function
    'Decide which probes pass the TM filter.
    'Arguments
    '    TM: An array of TMs
    '    TMFPars: An instance of the parameter class CTMFfilterPars.
    '    Filter: The return array of filter output values.
    On Error GoTo E

    Dim PSNames$           'name of current probeset
    Dim NumProbes          'number of probes in the recordset
    Dim Pos()               'Pos column from database
    Dim PosFilter() As Boolean
    Dim SeqLength          'Pos Filter column to database
                           'Length of the originating sequence.

    'Determine the number of probes.
    NumProbes = NumRecords(RSProbes)
    If NumProbes = 0 Then GoTo UpdateOnly

    'Start it up.
    PSName = RSProbes("PSName")
    Progress.ShowProgress "Calculating Pos Filter for Probeset " & PSName, 0,
NumProbes

    'Create space for database fields.
    ReDim Pos(0 To NumProbes - 1)
    ReDim PosFilter(0 To NumProbes - 1)

    'Get the Positions.
    GetField RSProbes, "Position", Pos

```

```

Public Sub CalcGenericFilter(Values, FilterPars As Object, Filter() As Boolean)
'-----'
'Function
'Decide which values pass the filter.
'Arguments
'  Values: An array of values to be filtered against.
'  FilterPars: The filter parameters
'  Filter: The returned filter settings.
'Notes
'  This routine is used to implement the common filter methods. FilterPars is
'  declared as an object rather than a specific class, and thus needs to have
'  only the members accessed on the execution path.
'  FilterPars must have data member Method.
'  If method is None, no other members are required.
'  If method is Min, then member Min is required.
'  If method is Max, then member Max is required.
'  If method is Distance, then members Max and Min are required.
'  If method is Range, then members Target and Distance are required.
'  If method is NBest, then members Target and NBest are required.
'  If method is Percent, then members Target and Percent are required.
'History
'  31-Jul-97: PW
'-----'

Dim Distance()           'absolute difference between values and target
Dim Target()              'target value
Dim MinValue#, MaxValue#
Dim Indexes()             'min and max of value
Dim NumValues#             'index array to track sort
                           'number of values we are working with
Dim Index#                'index
Dim Ns                      'number to set.

' Determine the number of probes we are calculating filter for.
NumValues = UBound(Values) + 1

'If method is "None", set all to true.
If FilterPars.Method = "None" Then
  For T = 0 To NumValues - 1
    Filter(T) = True
  Next T
  Exit Sub
End If

'If method is "Min", set all >= min to true.
If FilterPars.Method = "Min" Then
  For T = 0 To NumValues - 1
    Filter(T) = (Values(T) >= Val(FilterPars.Min))
  Next T
  Exit Sub
End If

'If method is "Max", set all <= max to true.
If FilterPars.Method = "Max" Then
  For T = 0 To NumValues - 1
    Filter(T) = (Values(T) <= Val(FilterPars.Max))
  Next T
  Exit Sub
End If

```

```

'If method is "InRange", set all >= min and <= max to true.
If FilterPars.Method = "InRange" Then
  For T = 0 To NumValues - 1
    Filter(T) = ((Values(T) >= Val(FilterPars.Min)) And (Values(T) <
= Val(FilterPars.Max)))
  Next T
  Exit Sub
End If

'For other methods, we have to do more work!
'Size the arrays
ReDim Distance(NumValues - 1)
ReDim Index(NumValues - 1)

'Find the minimum and maximum values.
For T = 0 To NumValues - 1
  If minValue > Values(T) Then minValue = Values(T)
  If maxValue < Values(T) Then maxValue = Values(T)
Next T

'Choose the target value.
If (FilterPars.Method = "Distance") Or -
(FilterPars.Method = "NBest") Or -
(FilterPars.Method = "Percent") Or -
Then Target = FilterPars.Target
If (FilterPars.Method = "NLowest") Or -
(FilterPars.Method = "PercentLowest") -
Then Target = minValue
If (FilterPars.Method = "NHighest") Or -
(FilterPars.Method = "PercentHighest") -
Then Target = maxValue

'Compute distances of values from target; set up index and filter.
For T = 0 To NumValues - 1
  Distance(T) = Abs(Values(T) - Target)
  Index(T) = T
  Filter(T) = False
Next T

'If method is "Distance", set all elements within distance as true.
If FilterPars.Method = "Distance" Then
  For T = 0 To NumValues - 1
    If Distance(T) = Distance(Index(T)) Then
      Filter(T) = True
    End If
  Next T
  Exit Sub
End If

'Choose the number to set.
If (FilterPars.Method = "NBest") Or -
(FilterPars.Method = "NLowest") Or -
(FilterPars.Method = "NHighest") -
Then N = FilterPars.NBest
If (FilterPars.Method = "Percent") Or -
(FilterPars.Method = "PercentLowest") Or -
(FilterPars.Method = "PercentHighest") -
Then N = NumValues * FilterPars.Percent / 100 - 1

```

```

If N >= NumValues Then N = NumValues - 1
' Sort on distance from target.
Quicksort Distance, Indxx, 0, NumValues - 1
'Set the best.
For T = 0 To N - 1
    Filter(Index(T)) = True
Next T
End Sub

Public sub CalcRunFilter(Run&(), RunFPars As CRUnFilterPars, Filter() As
Boolean)
' Function
' Decide which probes pass the Run filter.
' Arguments
' Run: An array of Runs
' RunFPars: An instance of the parameter class CRUnFilterPars.
' Filter: The return array of filter output values.
Dim OldMin
' The special method "All" can be handled by using the method Min, with min=1.
If RunFPars.Method = "All" Then
    OldMin = RunFPars.Min
    RunFPars.Min = 1
    RunFPars.Method = "Min"
    CalcGenericFilter Run, RunFPars, Filter
    RunFPars.Method = "All"
    RunFPars.Min = OldMin
    Exit Sub
End If
'otherwise, filters are generic.
CalcGenericFilter Run, RunFPars, Filter
End Sub

Public Sub CalcdGMFilter(dGM&(), dGMPars As cdGMFilterPars, Filter() As
Boolean)
' Function
' Decide which probes pass the dGM filter.
' Arguments
' dGM: An array of dGMs
' dGMPars: An instance of the parameter class cdGMFilterPars.
' Filter: The return array of filter output values.
CalcGenericFilter dGM, dGMPars, Filter
End Sub

Public Sub CalchomoFilter(Homology&(), HPPars As chmOFilterPars, Filter() As
Boolean)
' Function
' Decide which probes pass the homology filter.
' Arguments
' Homology: An array of homologies
' HPPars: An instance of the parameter class chmOFilterPars.
' Filter: The return array of filter output values.
CalcGenericFilter Homology, HPPars, Filter
End Sub

```

```

End Sub

'Grab the data.
EntrezData = frmEntrezGrab.Entrez.GetGenBankData(Accession, "")

'Create CRIFs out of LFS.
For 1 = 1 To Len(EntrezData)
  If (Mid$(EntrezData, 1, 1) = vbLf) Then EntrezData2 = EntrezData2 + vbCR
  EntrezData2 = EntrezData2 + Mid$(EntrezData, 1, 1)
Next 1

'Write the acquired data out to a file.
GBFfileName = frmMain.cdgSeqs.InitDir & "\\" & Accession & ".gb"
Open GBFfileName For Output As #1
Print #1, EntrezData2
Close #1

'Load the database.
SeqLoadDB ReaderRecord(GBFfileName), RS

'Remove the file.
If Not KeepFile Then Kill GBFfileName

Exit Sub
E: Debug.Print "Error in EntrezLoadDB"
Err.Raise Err.Number, , Err.Description
End Sub

Public Sub SeqLoadDB (SeqRC As SeqRecord, RS As Recordset)

'Function
'  Create a new record in the Sequence table, populate from SeqRC
'  Arguments
'    SeqRC: A sequence structure, as read from file, etc.
'    RS: The recordset into which the sequence should be loaded.
'  Errors:
'    Attempting to create a new sequence record whose Accession matches a
'    previously inserted sequence causes a DB error.

On Error GoTo E

BeginTrans
AllOnDbgClick = False
RS.AddNew
RS.Fields("Header") = SeqRC.Header
RS.Fields("Accession") = SeqRC.Accession
RS.Fields("Locus") = SeqRC.Locus
RS.Fields("Length") = SeqRC.Length
RS.Fields("Sequence") = SeqRC.Sequence
RS.Update
RS.Bookmark = RS.LastModified
CommitTrans
DoEvents
AllOnDbgClick = True
Exit Sub
E: Debug.Print "Error in SeqLoadDB"
Rollback
DoEvents

Attribute VB_Name = "GenBank"
Option Explicit

Private Const Gterminators = "/" 'record terminator in Genbank files
Private Const MaxGBFileLength = 100000 'maximum allowed Genbank file length

Public Sub EntrezLoadDB(Accessions$, RS As Recordset, KeepFile As Boolean)
'Function
'  Grab a sequence using the Entrez CGI.
'  Accession: The accession of the required sequence.
'  RS: The recordset into which this sequence should be loaded.
'  KeepFile: Controls whether the downloaded file is retained or removed.

On Error GoTo E

Dim GBfileName$ 'Genbank sequence file for data.
Dim EntrezData$, EntrezData2$ 'GB Entries.
Dim i$
```

```

AllowdbgClick = True
Err.Raise Err.Number, , Err.Description
End Sub

Public Function ReadGBRecord(FileName$) As SeqRecord
'-----
'Function
'  Read a GenBank record from a file into a seqRecord.
'Arguments
'  FileName: The filename of the GB record.
'Returns
'  The GenBank record as a seqRecord.
'Errors:
'  Returned to caller.
'  1. Check file is less than MaxGBFileLength
'  2. Check that all required fields are present.
'On Error GoTo E

Dim GBFile$          'GenBank file
Dim FileLength$      'file length (chars)
Dim LineLength$      'length of current line (chars)

Dim GBText$          'entire record
Dim GBItem$          'one line
Dim Keyfield$        'item keyfield
Dim foos$            'bit bucket

Dim GB As SeqRecord  'the returned record

'open the file
GBFile = FreeFile
Open FileName For Binary As #GBFile
Loop

'read in the file as one long string
FileLength = FileLen(FileName)
If FileLength > MaxGBFileLength Then Err.Raise pErrGBFileLength
GBText = Input(FileLength, #GBFile)

'check for required fields
If Instr(GBText, "LOCUS") = 0 Then Err.Raise pErrGBFileFormat, , "No LOCUS
field found in GB file."
If Instr(GBText, "ACCESSION") = 0 Then Err.Raise pErrGBFileFormat, , "No
ACCESSION field found in GB file."
If Instr(GBText, GBTerminator) = 0 Then Err.Raise pErrGBFileFormat, , "No
terminator (/) found in GB file."
'process items
Do While True

'extract an item
LineLength = Instr(GBText, vbCrLf) - 1
GBItem = Left(GBText, LineLength)
GB.Header = GB.Header & GBItem & vbCrLf
GBText = Right(GBText, FileLength - LineLength - 2)
FileLength = FileLength - LineLength - 2

If Left(GBItem, 2) = GBTerminator Then

```

```

    Goto breakwhile1
End If

'extract keyfield, and process item
Keyfield = StrField(GBItem, LineLength, 10)
'if keyfield is not numeric -> header
If IsNumeric(Keyfield) = False Then
  'if keyfield occupies first column -> keyword
  If Left(Keyfield, 1) <> " " Then
    'process keywords
    Select Case Keyfield
    Case "LOCUS"
      foo = StrField(GBItem, LineLength, 2)
      GB.Locus = StrField(GBItem, LineLength, 10)
    Case "ACCESSION"
      foo = StrField(GBItem, LineLength, 2)
      GB.Accession = StrField(GBItem, LineLength, 6)
    End Select
    'if keyfield has only two blanks -> subkeyword
    ElseIf Left(Keyfield, 3) <> " " Then
      'if keyfield has only 4 blanks -> feature code
      ElseIf Left(Keyfield, 5) <> " " Then
        'it must be a continuation line
        Else
          End If
        Else
          'sequence has started
          Do While LineLength > 10
            GB.Sequence = GB.Sequence & Left(GBItem, 10)
            GBItem = Right(GBItem, LineLength - 11)
            LineLength = LineLength - 11
          Loop
          If LineLength > 0 Then
            GB.Sequence = GB.Sequence & GBItem
            End If
          End If
        End If
      End If
    End If
  End If

  'strip any whitespace.
  GB.Sequence = PackSequence(GB.Sequence)

  'Check sequence length.
  If GB.Length <> Len(GB.Sequence) Then
    MsgBox "Sequence length differs from header specification."
    GB.Length = Len(GB.Sequence)
  End If

  ReadGBRecord = GB
  Exit Function
E: Debug.Print "Error in ReadGBRecord"
Err.Raise Err.Number, , Err.Description
End Function

```

```

Public Function ReadFASTAREcord(FASTAfile$) As SeqRecord
    'Function
    '  Read a FASTA record from a file into a SeqRecord.
    '  Arguments
    '    FASTA: The file descriptor for the FASTA file.
    '  Returns:
    '    The FASTA record as a SeqRecord.
    '  Errors:
    '    Returned to caller.
    '  Notes:
    '    The FASTA file must already be opened for input, and should be positioned
    '    at the start of a FASTA header. It will read an entire sequence, leaving
    '    the file positioned on the subsequent header.

On Error Goto E

Dim LineTexts$           'One line of the file
Dim SR As SeqRecord      'the returned record

'Read the header line, check it is legal.
Line Input #FASTAfile, LineText
If Left(LineText, 1) <> ">" Then Err.Raise pErrFASTAFormat, , "Header line does
not begin w/ >."

'Extract the length.
LineText = Right(LineText, Len(LineText) - InStr(LineText, "len") - 3)
If InStr(LineText, "#") <> 0 Then LineText = Left(LineText, InStr(LineText, "#") - 1)
SR.Length = CInt(LineText)

'Read sequence lines till we are done.
Do While (Len(SR.Sequence) <> SR.Length)
    Line Input #FASTAfile, LineText
    SR.Sequence = SR.Sequence & LineText
Loop

ReadFASTAREcord = SR

Exit Function
E: Debug.Print "Error in ReadFASTAREcord"
Err.Raise Err.Number, , Err.Description
End Function

Public Function ReadFASTAHeader(FASTAfile$) As SeqRecord
    'Function
    '  Read a FASTA header from a file into a SeqRecord.
    '  Arguments
    '    FASTA: The file descriptor for the FASTA header file.
    '  Returns:
    '    The FASTA record as a SeqRecord.
    '  Errors:
    '    Returned to caller.
    '  Notes:
    '    The FASTA header file must already be opened for input, and should be
    '    positioned at the start of a FASTA header.

```

```

On Error Goto E

Dim LineTexts      'one line of the file
Dim LineText$      'line processed to find length
Dim AccText$       'line processed to find accession
Dim SR As SeqRecord 'the returned record

'Read the header line, check it is legal.
Line Input #FASTAFile, LineText
If Left(LineText, 1) <> ">" Then Err.Raise PPErrMsgFormat, , "Header line does
not begin w/ >"

'Extract the accession.
LineText = Right(LineText, Len(LineText) - InStr(LineText, ">") - 3)
If InStr(LineText, ">") <> 0 Then LenText = Left(LineText, InStr(LineText, ">") - 1)
SR.Length = CInt(LineText)

'Extract the accession.
AccText = Right(LineText, Len(LineText) - InStr(LineText, "#gb=") - 2)
If InStr(AccText, "#") <> 0 Then AccText = Left(AccText, InStr(AccText, "#") - 1)
SR.Accession = LTrim(RTrim(AccText))

ReadFASTAHeader = SR

Exit Function
E: Debug.Print "Error in ReadFASTARecord"
Err.Raise Err.Number, , Err.Description
End Function

Public Function GBCookSeq(ByVal RawSeq$) As String
'Convert raw sequence (no spaces, etc) into more
'palatable form, with spaces every 10 nucleotides
'and newlines every 60 nucleotides.

Dim Bases$          'count the bases
Dim LineLength$     'remaining line length
LineLength = Len(RawSeq)
Bases = 1
Do While LineLength > 60
    GBCookSeq = GBCookSeq & StrField(RawSeq, LineLength, 10) & " "
    Format(Format(Bases, "-#####"), "000000") & " "
    Bases = Bases + 60
    Dim i6
    For i = 1 To 6
        GBCookSeq = GBCookSeq & StrField(RawSeq, LineLength, 10) & " "
    Next i
    GBCookSeq = GBCookSeq & vbCrLf
Loop
If Len(RawSeq) <> 0 Then
    GBCookSeq = GBCookSeq & StrField(RawSeq, LineLength, 10) & " "
    Format(Format(Bases, "-#####"), "000000") & " "
    Do While Len(RawSeq) > 10
        GBCookSeq = GBCookSeq & StrField(RawSeq, LineLength, 10) & " "
    Loop
End If
End Function

```

```

Loop
    GBCookSeq = GBCookSeq & RawSeq
End If
End Function

Public Function RTFGBCookSeq($RawSeq$) As String
    'Convert raw sequence (no spaces, etc) into more
    'palatable form, with spaces every 10 nucleotides
    'and newlines every 60 nucleotides.

    Dim Bases$&
    Dim LineLength$&
    'count the bases
    LineLength = Len($RawSeq$)
    Bases = 1
    Do While LineLength > 60
        RTFGBCookSeq = RTFGBCookSeq & $RawSeq$(1 To 60) & $<br>
        Format(Format(Bases, "####"), "00000") & $<br>
        Bases = Bases + 60
    Loop
    LineLength = Len($RawSeq$)
    For 1 = 1 To 6
        RTFGBCookSeq = RTFGBCookSeq & StrField($RawSeq$, LineLength, 10) & $<br>
    Next 1
    RTFGBCookSeq = RTFGBCookSeq & "\par"
    If Len($RawSeq$) <> 0 Then
        RTFGBCookSeq = RTFGBCookSeq & Format(Format(Bases, "####"), "00000") & $<br>
        Do While Len($RawSeq$) > 10
            RTFGBCookSeq = RTFGBCookSeq & StrField($RawSeq$, LineLength, 10) & $<br>
        Loop
        RTFGBCookSeq = RTFGBCookSeq & $RawSeq$
    End If
End Function

Attribute VB_Name = "SQL"
Option Explicit

Public Function BuildProbePSQuery($PName$)
    'Function
    'Build an SQL Query to select Probes from the Probes table
    'that pass all filters, and come from the named ProbeSet.
    'Arguments
    'Accession: The accession to search on.
    'PName: The ProbeSet name to search on.
    'Notes
    ' 2. All fields corresponding to a column (that is, all fields
    '     named in the Columns table) are returned, to match the layout
    '     of the Probes grid. The visibility of various columns is
    '     controlled by the column Visible properties.
    ' 3. The currently set filters are used as a further selection
    '     on the probes returned.
    '-----'
    Dim SelectC$, FromC$, FilterCS$, WhereCS$, OrderByCS
    Dim C$, F$&

    'Set up the select clause -- return all columns.
    SelectC = "SELECT *"
    'For C = 1 To PSColumns.Count
    '    If C <> 1 Then SelectC = SelectC & ","
    '    SelectC = SelectC & "[" & PSColumns(C).Name & "]"
    'Next C

    'Set up the the from clause -- always the Probes table.
    FromC = "FROM Probes"
    'Set up the where clause
    For F = 1 To PSColumns.Count
        If (PSColumns(F).IsFilter = True) Then
            If FilterC = "" Then
                FilterC = "([& PSColumns(F).Name & ] = True)"
            Else
                FilterC = FilterC & " AND ([& PSColumns(F).Name & ] = True)"
            End If
        End If
    Next F
    WhereC = "WHERE ( $PName$ = [& PSColumns(F).Name & ] )"
    If PSColumns("User Filter").IsFilter Then
        If FilterC <> "" Then
            If FilterC = "" Then
                WhereC = WhereC & " AND ( ([User Filter] = 2) OR ( ([User Filter] = 1)
                    AND [& FilterC & ]) ) "
            Else
                WhereC = WhereC & " AND ([User Filter] <> 0) ) "
            End If
        Else
            If FilterC <> "" Then
                WhereC = WhereC & " AND ( [User Filter] <> 0 ) "
            Else
                WhereC = WhereC & " "
            End If
        End If
    End If

    'Put it all together.
    BuildProbePSQuery = SelectC & FromC & WhereC & ";"
End Function

Public Function BuildProbePSQuery()
    'Function
    'Build an SQL Query to select Probes from the Probes table.
    'Arguments
    'Accession: The accession to search on.
    'PName: The ProbeSet name to search on.
    'Notes
    ' 1. Accession or PName = <none> is allowed, and results
    '     in an empty recordset.
    ' 2. All fields corresponding to a column (that is, all fields
    '     named in the Columns table) are returned, to match the layout
    '     of the Probes grid. The visibility of various columns is
    '     controlled by the column Visible properties.
    ' 3. The currently set filters are used as a further selection
    '     on the probes returned.
    '-----'
    Dim SelectC$, FromC$, FilterCS$, WhereCS$, OrderByCS
    Dim C$, F$&

```

```

'-- SelectC$, FromC$, FilterC$, WhereC$, OrderByC$          Option Explicit
Dim SelectC$, FromC$, FilterC$, WhereC$, OrderByC$          Public Type StatRecord
Dim C$, F$                                         Count As Long
Dim Col As column                                     Min As Double
                                             Median As Double
                                             Ninety As Double
                                             Max As Double
                                             Range As Double
                                             Average As Double
                                             Std As Double
End Type

Public Function Statistics(Vector() As StatRecord
'Function
'Calculate all statistics of a vector.
'
Dim L$, U$                                         Dim L$, U$, Min$, Max$, Total$, Total12$
Dim Index$()                                       Dim Index$()
Dim I$                                           Dim I$ To U$ - L$ + 1
ReDim Index(I$ To U$)
Dim I$                                           Next I$ To U$ - L$ + 1
Min$ = Vector(0)                                     Total$ = Total$ + Vector(I$) * Statistics.Count
Max$ = Vector(0)                                     If Vector(I$) > Max$ Then Max$ = Vector(I$)
Range$ = Vector(0)                                    If Vector(I$) < Min$ Then Min$ = Vector(I$)
Statistics.Average = Total$ / Statistics.Count
Statistics.Std = Sqr((Total12 / Statistics.Count) - Statistics.Average * Statistics.Average)

'Sort.
QuickSort Vector, Index, L$, U$

'Percentiles.
Statistics.Ten = Vector(1 + (Statistics.Count - 1) * 0.1)
Statistics.Median = Vector(1 + (Statistics.Count - 1) * 0.5)
Statistics.Ninety = Vector(1 + (Statistics.Count - 1) * 0.9)

Attribute VB_Name = "Stats"

```

```

'-- SelectC$, FromC$, FilterC$, WhereC$, OrderByC$          Option Explicit
Dim SelectC$, FromC$, FilterC$, WhereC$, OrderByC$          Public Type StatRecord
Dim C$, F$                                         Count As Long
Dim Col As column                                     Min As Double
                                             Median As Double
                                             Ninety As Double
                                             Max As Double
                                             Range As Double
                                             Average As Double
                                             Std As Double
End Type

Public Function Statistics(Vector() As StatRecord
'Function
'Calculate all statistics of a vector.
'
Dim L$, U$                                         Dim L$, U$, Min$, Max$, Total$, Total12$
Dim Index$()                                       Dim Index$()
Dim I$                                           Dim I$ To U$ - L$ + 1
ReDim Index(I$ To U$)
Dim I$                                           Next I$ To U$ - L$ + 1
Min$ = Vector(0)                                     Total$ = Total$ + Vector(I$) * Statistics.Count
Max$ = Vector(0)                                     If Vector(I$) > Max$ Then Max$ = Vector(I$)
Range$ = Vector(0)                                    If Vector(I$) < Min$ Then Min$ = Vector(I$)
Statistics.Average = Total$ / Statistics.Count
Statistics.Std = Sqr((Total12 / Statistics.Count) - Statistics.Average * Statistics.Average)

'Sort.
QuickSort Vector, Index, L$, U$

'Percentiles.
Statistics.Ten = Vector(1 + (Statistics.Count - 1) * 0.1)
Statistics.Median = Vector(1 + (Statistics.Count - 1) * 0.5)
Statistics.Ninety = Vector(1 + (Statistics.Count - 1) * 0.9)

Attribute VB_Name = "Stats"

```

```

'-- Set up the select clause -- return all columns.
SelectC = "SELECT *"
For C = 1 To PSColumns.Count
  If C > 1 Then SelectC = SelectC & ","
  SelectC = SelectC & "Probes.[*] & PSColumns(C).Name & "]
Next C

'-- Set up the from clause -- from the SelectProbes query.
FromC = "FROM SelectProbes"

'-- Set up the where clause -- filter the rows.
'For F = 1 To PSColumns.Count
'  For Each Col In PSColumns
'    If (Col.IsFilter = True) Then
'      If FilterC = "" Then
'        FilterC = "[*] & Col.Name & " = True)"
'      Else
'        FilterC = FilterC & " AND ([*] & col.Name & "] = True)"
'      End If
'    End If
'  Next Col
WhereC = "WHERE ( Selected = True )"
If PSColumns("User Filter").IsFilter Then
  If FilterC <> "" Then
    WhereC = WhereC & " AND ( [User Filter] = 2 ) OR ( [User Filter] = 1 )"
    AND " & FilterC & ")"
  Else
    WhereC = WhereC & " AND ([User Filter] <> 0) )"
  End If
Else
  If FilterC <> "" Then
    WhereC = WhereC & " AND ( [*] & FilterC & " ) )"
  Else
    WhereC = WhereC & " )"
  End If
End If

'-- Set up the order by clause.
'Default to sorting by position, then check columns.
OrderByC = "Position"
For Each Col In PSColumns
  If (Col.Sort = "Ascending") Then OrderByC = "[*] & Col.Name & " ASC "
  If (Col.Sort = "Descending") Then OrderByC = "[*] & Col.Name & " DESC "
Next Col
OrderByC = "ORDER BY Probes.Accession, Probes.PSName, " & OrderByC

'Put it all together.
BuildProbesQuery = SelectC & FromC & WhereC & OrderByC & ";"

End Function

Attribute VB_Name = "Stats"

```

```

Private RNAHPP_Stack#(0 To 3, 0 To 3, 0 To 3, 0 To 3) RNAHPP_Stack
Private RNAHPP_TStack#(0 To 3, 0 To 3, 0 To 3, 0 To 3) RNAHPP_TStack
Private RNAHPP_Dangle#(0 To 1, 0 To 3, 0 To 3, 0 To 3) RNAHPP_Dangle
Private RNAHPP_Dangle#(0 To 2, 0 To 3, 0 To 3, 0 To 3) RNAHPP_Dangle
Private RNAHPP_TLoop#(0 To NumTetraloops - 1) RNAHPP_TLoop
Private RNAHPP_TLoop#(0 To NumTetraloops - 1) RNAHPP_TLoop

Attribute VB_Name = "Thermo"
Option Explicit

'Where is no good reason to have two sets of thermodynamic parameters,
'other than history...
Parameters used by TM calculations.
Public Const RGasf = 1.987

Private DNA_Duplex#(0 To 3, 0 To 3) DNA_Duplex
Private DNA_Duplex#(0 To 3, 0 To 3) DNA_Duplex
Private DNA_InitATScf; DNA_InitATScf
Private DNA_SelfScf; DNA_SelfScf
Private DNA_EndTHf; DNA_EndTHf
Private DR_Duplex#(0 To 3, 0 To 3) DR_Duplex
Private DR_Duplex#(0 To 3, 0 To 3) DR_Duplex
Private DR_3InitHf; DR_3InitHf
Private DR_3InitHf; DR_3InitHf
Private DR_InitHf; DR_InitHf
Private DR_InitHf; DR_InitHf

'Gas constant, cal/mol/deg K.
'DNA/DNA nearest neighbor enthalpies.
'DNA/RNA nearest neighbor entropies.
'DNA/DNA initiation entropies.
'DNA/DNA self-symmetry entropies.
'DNA/DNA end enthalpy.
'DNA/RNA nearest neighbor enthalpies.
'DNA/RNA nearest neighbor entropies.
'DNA/RNA initiation enthalpy/entropy.

Parameters used by hairpin calculations.
'Zuker provides H and G @ 37. So on load, calculate S, then recalculate G as
needed.
Private Const NumZukerLoopsf = 30
'Number of special case loops.
'Number of special case loop lengths.
'The special case loop lengths.

Private Const RNA_2MiscLoopf = 1.079
'Constants for larger-loop
Private Const DNA_2MiscLoopf = 1.079
'Minimum allowed loop.
'Number of special tetraloops.
'Number of special tetraloops.
'Indices of tetraloops.

'DNA free energy.
Private DNAHPP_Stack#(0 To 3, 0 To 3, 0 To 3, 0 To 3) DNAHPP_Stack
Private DNAHPP_TStack#(0 To 3, 0 To 3, 0 To 3, 0 To 3) DNAHPP_TStack
Private DNAHPP_Dangle#(0 To 1, 0 To 3, 0 To 3, 0 To 3) DNAHPP_Dangle
Private DNAHPP_Dangle#(0 To 2, 0 To 3, 0 To 3, 0 To 3) DNAHPP_Dangle
Private DNAHPP_TLoop#(0 To NumTetraloops - 1) DNAHPP_TLoop
Private DNAHPP_TLoop#(0 To NumTetraloops - 1) DNAHPP_TLoop

'DNA enthalpy.
Private DNAHPP_Stacks#(0 To 3, 0 To 3, 0 To 3, 0 To 3) DNAHPP_Stacks
Private DNAHPP_TStacks#(0 To 3, 0 To 3, 0 To 3, 0 To 3) DNAHPP_TStacks
Private DNAHPP_Dangles#(0 To 1, 0 To 3, 0 To 3, 0 To 3) DNAHPP_Dangles
Private DNAHPP_Dangles#(0 To 2, 0 To 3, 0 To 3, 0 To 3) DNAHPP_Dangles
Private DNAHPP_TLoops#(0 To NumTetraloops - 1) DNAHPP_TLoops
Private DNAHPP_TLoops#(0 To NumTetraloops - 1) DNAHPP_TLoops

'RNA enthalpy.
Private RNAHPP_Stacks#(0 To 3, 0 To 3, 0 To 3, 0 To 3) RNAHPP_Stacks
Private RNAHPP_TStacks#(0 To 3, 0 To 3, 0 To 3, 0 To 3) RNAHPP_TStacks
Private RNAHPP_Dangles#(0 To 1, 0 To 3, 0 To 3, 0 To 3) RNAHPP_Dangles
Private RNAHPP_Dangles#(0 To 2, 0 To 3, 0 To 3, 0 To 3) RNAHPP_Dangles
Private RNAHPP_TLoops#(0 To NumTetraloops - 1) RNAHPP_TLoops
Private RNAHPP_TLoops#(0 To NumTetraloops - 1) RNAHPP_TLoops

'RNA entropy.
Private RNAHPP_Stacks#(0 To 3, 0 To 3, 0 To 3, 0 To 3) RNAHPP_Stacks
Private RNAHPP_TStacks#(0 To 3, 0 To 3, 0 To 3, 0 To 3) RNAHPP_TStacks
Private RNAHPP_Dangles#(0 To 1, 0 To 3, 0 To 3, 0 To 3) RNAHPP_Dangles
Private RNAHPP_Dangles#(0 To 2, 0 To 3, 0 To 3, 0 To 3) RNAHPP_Dangles
Private RNAHPP_TLoops#(0 To NumTetraloops - 1) RNAHPP_TLoops
Private RNAHPP_TLoops#(0 To NumTetraloops - 1) RNAHPP_TLoops

'Function
'Calculate S given G, H, and T, for all elements of array S.
Arguments
'Dims: The number of dimensions of G, H, and S.
'G: The given free energies.
'H: The given enthalpies.
'T: The temperature at which free energy was calculated.
'S: The returned entropies.
Notes
1. The reason for this functions existence is that Zuker provides
G at 37, and H. S is needed to change temperatures.
Dim I6, J6, K6, L6
T = T + 273.15
Select Case Dims
Case 1
For I = LBound(G) To UBound(G)
S(I) = (G(I) - H(I)) / T
Next I
Case 2
For I = LBound(G, 1) To UBound(G, 1)
For J = LBound(G, 2) To UBound(G, 2)
S(I, J) = (G(I, J) - H(I, J)) / T
Next I
Case 4
For I = LBound(G, 1) To UBound(G, 1)
For J = LBound(G, 2) To UBound(G, 2)
For K = LBound(G, 3) To UBound(G, 3)
For L = LBound(G, 4) To UBound(G, 4)
S(I, J, K, L) = (G(I, J, K, L) - H(I, J, K, L)) / T
Next I
Case 5
For I = LBound(G, 1) To UBound(G, 1)
For J = LBound(G, 2) To UBound(G, 2)
For K = LBound(G, 3) To UBound(G, 3)
For L = LBound(G, 4) To UBound(G, 4)
S(I, J, K, L) = (G(I, J, K, L) - H(I, J, K, L)) / T
Next I
Next K
Next L
Next J
Next I
End Select

```

```

End Sub

    Private sub CalcGFromHS(Dims$, H(), S(), ByVal T$, G())
        'Function
        'Calculate G given H, S, and T, for all elements of array G.
        'Arguments
        ' Dims: The number of dimensions of G, H, and S.
        ' H: The given enthalpies.
        ' S: The given entropies.
        ' T: The temperature at which free energy is required.
        ' G: The returned free energies.
    End Sub

    Dim I$, J$, K$, L$
    T = T + 273.15
    Select Case Dims
    Case 1
        For I = LBound(G) To UBound(G)
            G(I) = 0.1 * CDBL(CINT(10$ * (H(I) + S(I) * T)))
        Next I
    Case 2
        For I = LBound(G, 1) To UBound(G, 1)
            For J = LBound(G, 2) To UBound(G, 2)
                For K = LBound(G, 3) To UBound(G, 3)
                    For L = LBound(G, 4) To UBound(G, 4)
                        G(I, J, K, L) = 0.1 * CDBL(CINT(10$ * (H(I, J) + S(I, J) * T)))
                    Next L
                Next K
            Next J
        Next I
    End Select
    End Sub

    Private Function DNA_BestHairpin$(seq$())
        'Function
        'Compute the most stable hairpin for the given sequence.
        'Arguments
        ' seq: The numerical representation of the sequence.
    End Function

    Dim Thishairpin#
    Dim NumBases#
    Dim FP$, TP$
    DNA_BestHairpin = UBound(seq) - LBound(seq) + 1
    For FP = 0 To NumBases - 2MinLoop - 1
        For TP = FP + 2MinLoop + 1 To NumBases - 1
            Thishairpin = RNA_Hairpin(Seq, FP, TP)
            If Thishairpin < DNA_BestHairpin Then DNA_BestHairpin = Thishairpin
        Next TP
    Next FP
    End Function

    Private Function RNA_BestHairpin$(seq$())
        'Function
        'Compute the most stable hairpin for the given sequence.
        'Arguments
        ' seq: The numerical representation of the sequence.
    End Function

    Dim Thishairpin#
    Dim NumBases#
    Dim FP$, TP$
    RNA_BestHairpin = 1000#
    NumBases = UBound(seq) - LBound(seq) + 1
    For FP = 0 To NumBases - 2MinLoop - 1
        For TP = FP + 2MinLoop + 1 To NumBases - 1
            Thishairpin = RNA_Hairpin(Seq, FP, TP)
            If Thishairpin < RNA_BestHairpin Then RNA_BestHairpin = Thishairpin
        Next TP
    Next FP
    End Function

    Private Function RNA_Hairpin$(seq$(), ByVal FP$, ByVal TP$)
        'Function
        'Calculate the free energy of a RNA hairpin
        'Arguments
        ' seq: The sequence, in numeric representation.
        ' FP: The index of the 5'-end base that closes the loop.
        ' TP: The index of the 3' end base that closes the loop.
    End Function

    Dim Energies$()
    Dim NumBases, StemLength, S#
    'Check whether this pair can close the loop.
    RNA_Hairpin = 1000#
    Select Case Seq(FP)
    Case 0
        If Seq(TP) <> 3 Then Exit Function
    Case 1
        If Seq(TP) <> 2 Then Exit Function
        If Seq(TP) <> 1 And Seq(TP) >> 3 Then Exit Function
    Case 2
        If (Seq(TP) <> 1 And Seq(TP) >> 3) Then Exit Function
    Case 3
        If (Seq(TP) <> 0 And Seq(TP) >> 2) Then Exit Function
    End Select
    'Calculate sequence size.
    NumBases = UBound(seq) - LBound(seq) + 1
    'setup to store all possible energies.

```

```

Redim Energies(UBound(Seq) - LBound(Seq) + 1)

' Start the energy calculations with the loop.
Energies(0) = RNAHP_Loop(2, TP - FP - 2);

' Add the stacking interaction of the first mismatch in the loop.
Energies(0) = Energies(0) + RNAHP_TStackG(seq(FP), Seq(FP + 1), seq(TP),
~ 11)

' Loop over stem members
S = 0
Do While True
    FP = FP -
    TP = TP +
    'Check that there are still bases to process.
    If (FP >= 0) And (TP < NumBases) Then
        'Check that we are still in the helix.
        Select Case Seq(FP)
        Case 0
            If Seq(TP) <> 3 Then Exit Do
        Case 1
            If seq(TP) >> 2 Then Exit Do
            If (seq(TP) <> 1 And seq(TP) <> 3) Then Exit Do
        Case 2
            If (seq(TP) <> 0 And seq(TP) <> 2) Then Exit Do
        Case 3
            If (Seq(TP) <> 0 And Seq(TP) <> 2) Then Exit Do
        End Select
        'Calculate sequence size.
        Numbases = UBound(Seq) - LBound(Seq) + 1
        'Setup to store all possible energies.
        Redim Energies(UBound(Seq) - LBound(Seq) + 1)
        'Start the energy calculations with the loop.
        Energies(0) = DNAHP_LoopG12(TP - FP - 2);
        'Add the stacking interaction of the first mismatch in the loop.
        Energies(0) = Energies(0) + DNAHP_TStackG(seq(FP), seq(TP), seq(TP,
~ 11))

        'Loop over stem members
        S = 0
        Do While True
            If Seq(TP) <> 3 Then Exit Do
            If (TP < NumBases) Then
                'Check that we are still in the helix.
                Select Case Seq(FP)
                Case 0
                    If Seq(TP) <> 3 Then Exit Do
                    If (TP >= 0) And (TP < NumBases) Then
                        'Find the minimum energy.
                        RNA_Hairpin = 1000#
                        StemLength = S
                        For S = 0 To StemLength
                            If Energies(S) < RNA_Hairpin Then RNA_Hairpin = Energies(S)
                        Next S
                End If
            End If
        End Do
    End If
Loop

'Add dangles, if they exist.
If (FP >= 0) Then Energies(S) = Energies(S) + RNAHP_DangleG(1, Seq(TP - 1),
Seq(FP + 1), Seq(FP))
If (TP < NumBases) Then Energies(S) = Energies(S) + RNAHP_DangleG(0, Seq(TP -
1), Seq(TP), Seq(FP + 1))

'Find the minimum energy.
RNA_Hairpin = 1000#
StemLength = S
For S = 0 To StemLength
    If Energies(S) < RNA_Hairpin Then RNA_Hairpin = Energies(S)
Next S

```

```

Case 3
  If (seq(TP) < 0 And Seq(TP) <> 2) Then Exit Do
  End Select

  'Add the next stacking term
  Energies(S + 1) = Energies(S) + DNAHP_StackG(Seq(TP), Seq(TP + 1),
Seq(TP), Seq(TP - 1))

  'Record the energy if the helix breaks here.
  Energies(S) = Energies(S) + DNAHP_DangleG(1, seq(TP - 1), seq(TP + 1),
Seq(TP)) + -DNAHP_DangleG(0, seq(TP - 1), seq(TP), seq(TP + 1))

  S = S + 1
  Else
    Exit Do
  End If
Loop

'Add dangles, if they exist.
If (TP >= 0) Then Energies(S) = Energies(S) + DNAHP_DangleG(1, seq(TP - 1),
Seq(TP + 1), Seq(TP))
If (TP < NumberBases) Then Energies(S) = Energies(S) + DNAHP_DangleG(0, seq(TP -
1), seq(TP), seq(TP + 1))

'Find the minimum energy.
DNA_Hairpin = 1000#
StemLength = S
For S = 0 To StemLength
  If Energies(S) < DNA_Hairpin Then DNA_Hairpin = Energies(S)
Next S
End Function

```

```

Public Sub InitThermoParams()
  DNA_InitGCs = -5.9
  DNA_InitialT = -9
  DNA_Selfs = -1.4

  'Initialize all the fixed parameters used for Thermo calculations.
  'Notes:
  1. These parameters are essentially constants, but are placed in
  arrays for ease-of-use. Thus this routine must be called by
  Main to initialize everything.
  2. The nearest-neighbor parameters are accessed by indexing by
  the first then second base, with A->0, C->1, G->2, T/U->3
  3. The hairpin parameters are loaded from files by this routine,
  and the entropy matrices filled in. After this operation, the
  free energy matrices may be overwritten at any time
  'files opened for Zuker parameters.
  Dim FileName$ 'files opened for Zuker parameters.
  Dim foot$ 'bitbucket.

  'Initialize the DNA Duplex enthalpy and entropy.
  'These parameters are from SantaLucia et al. Biochemistry, v. 35, pp 3555.
  'DR_DuplexH is in kcal/mol, Duplexes in cal/mol/deg K.

```

```

CalcsFromGH 1, RNAHP_TLoopG, DNAHP_TLoopH, 37#, RNAHP_TLoops

'GC
'GG
'GU
'UA
'UC
'TUG
'UU

DR_InitH = 1.9

DR_DuplexH(2, 1) = -8#
DR_DuplexH(2, 2) = -9.3
DR_DuplexH(2, 3) = -5.9
DR_DuplexH(3, 1) = -7.8
DR_DuplexH(3, 2) = -5.5
DR_DuplexH(3, 3) = -7.8

DR_DuplexS(0, 0) = -36.4
DR_DuplexS(0, 1) = -21.6
DR_DuplexS(0, 2) = -19.7
DR_DuplexS(0, 3) = -23.9
DR_DuplexS(1, 0) = -28.4
DR_DuplexS(1, 1) = -31.9
DR_DuplexS(1, 2) = -47.1
DR_DuplexS(1, 3) = -23.5
DR_DuplexS(2, 0) = -22.9
DR_DuplexS(2, 1) = -17.1
DR_DuplexS(2, 2) = -23.2
DR_DuplexS(2, 3) = -12.3
DR_DuplexS(3, 0) = -23.2
DR_DuplexS(3, 1) = -13.5
DR_DuplexS(3, 2) = -26.1
DR_DuplexS(3, 3) = -21.9

DR_InitS = -3.9

'Load the Zuker hairpin parameters.
FileName = App.Path & "\stack.dat.pw"
ReadZuker FileName, "Stack", DNAHP_StackH, foo
FileName = App.Path & "\stack.dhd.pw"
ReadZuker FileName, "Stack", DNAHP_StackH, foo
CalcsFromGH 4, DNAHP_TStackH, DNAHP_StackH, 37#, DNAHP_Stacks

FileName = App.Path & "\tstackh.dat.pw"
ReadZuker FileName, "Stack", DNAHP_TStackH, foo
FileName = App.Path & "\tstack.dhd.pw"
ReadZuker FileName, "Stack", DNAHP_TStackH, foo
CalcsFromGH 4, RNAHP_TStackH, RNAHP_StackH, 37#, RNAHP_Stacks

FileName = App.Path & "\tstack.dat.pw"
ReadZuker FileName, "Stack", RNAHP_TStackH, foo
FileName = App.Path & "\tstack.dhd.pw"
ReadZuker FileName, "Stack", RNAHP_TStackH, foo
CalcsFromGH 4, RNAHP_TStackH, RNAHP_StackH, 37#, RNAHP_TStacks

FileName = App.Path & "\dangle.dat.pw"
ReadZuker FileName, "Dangle", RNAHP_DangleH, foo
FileName = App.Path & "\dangle.dhd.pw"
ReadZuker FileName, "Dangle", RNAHP_DangleH, foo
CalcsFromGH 4, RNAHP_DangleH, RNAHP_DangleH, 37#, RNAHP_DangleH

FileName = App.Path & "\loop.dat.pw"
ReadZuker FileName, "Loop", RNAHP_LoopG, ZLoopLengths
FileName = App.Path & "\loop.dhd.pw"
ReadZuker FileName, "Loop", RNAHP_LoopH, ZLoopLengths
CalcsFromGH 4, RNAHP_LoopG, RNAHP_LoopH, 37#, RNAHP_Loops

FileName = App.Path & "\tloop.dat.pw"
ReadZuker FileName, "Tloop", RNAHP_TLoopG, ZTetraloops
FileName = App.Path & "\tloop.dhd.pw"
ReadZuker FileName, "Tloop", RNAHP_TLoopH, ZTetraloops
CalcsFromGH 1, RNAHP_TLoopG, RNAHP_TLoopH, 37#, RNAHP_TLoops

End Sub

Private Sub ReadZuker(FileNameS, ZTypeS, Param1(), FirstCol())
    'Function
    'Read thermodynamic parameters for structure calculations.
    'Arguments
    'Filename: The data file.
    'ZType: The type of parameter array (see below).
    'Param1: The parameter array to be filled in.
    'FirstCol: The values from the first column.
    'Notes
    '1. This routine reads data files originally designed to be
        MFC01D; the files must be edited to place a # sign on line
        do not contain data, and to replace all the "# entries".
    '2. There are several types of parameter files. Type "stack"
        gives the parameters for stacking one base pair over another.
        and thus has 256 entries. Type "dangle" gives the parameters
        for dangling a base over a pair, and thus has 64 parameters.
        The other types are particular to the parameters being
        presented.
    '3. The FirstCol argument is used only by Loop and Tetraloops
    '4. In all these files, the order of bases is ACCU->0,1,2,3
End Sub

```

```

Dim ZFile$           'parameter file
Dim FileLength$     'length of parameter file
Dim LineLength$      'length of one line
Dim WordLength$      'length of one word
Dim Filestr$          'text of read-in file
Dim Linestr$          'text of one line
Dim Wordstr$          'text of one word
Dim A$, B$, X$, Y$, L$, G$, T$ 'loop indices
Dim NumSeq(0 To 3)    'numeric representation of tetraloop sequences

'Open the file, read it in.
ZFile = Freefile
Open FileName For Binary As #ZFile
FileLength = Filelen(Filename)
Filestr = Input(FileLength, #ZFile)

'Choose how to parse the file.
Select Case ZType
Case "Stack"
'These entries represent stacking of one base pair over another:
    .5'-AX-3'
    .2'-BY-5'.
    'In each row, Y varies fast, B varies slow.
    'By row, X varies fast, and A varies slowly.
    'Index order used is param(A,X,B,Y).
    For A = 0 To 3
        For B = 0 To 3
            NextLine Filestr, FileLength, Linestr, LineLength, "##"
            For Y = 0 To 3
                NextWord Linestr, LineLength, Wordstr, WordLength
                Param(A, X, B, Y) = CDBl(Wordstr)
                Next Y
            Next B
        Next A
    Case "Dangle"
'These entries represent dangles of 3' ends, then the 5' ends:
    .5'-BX-3'
    .3'-B -5'
    .and
    .5'-A -3'
    .3'-BY-5'
    'In each row, X or Y varies fast, and B varies slowly.
    'By row, A varies.
    'Index order used is param(0,A,X,B) for 3', and param(1,A,B,Y) for the 5'.
    For A = 0 To 3
        NextLine Filestr, FileLength, Linestr, LineLength, "##"
        Param(0, A, X, B) = CDBl(Wordstr)
    Next X

```

```

    Next A
    For A = 0 To 3
        NextLine Filestr, FileLength, Linestr, LineLength, "##"
        For B = 0 To 3
            For Y = 0 To 3
                NextWord Linestr, LineLength, Wordstr, WordLength
                Param(1, A, B, Y) = CDBl(Wordstr)
            Next Y
        Next B
    Next A
    Case "Loop"
'These entries represent internal=0, bulge=1, and hairpin=2 loops.
    'By row, loop length L varies.
    'Index order used is param(looptype,L)
    'FirstCol is filled from the first column, representing loop lengths.
    For L = 0 To NumTetraloops - 1
        NextLine Filestr, FileLength, Linestr, LineLength, "##"
        FirstCol(L) = CLng(Wordstr)
        NextWord Linestr, LineLength, Wordstr, WordLength
        FirstCol(L) = CLng(Wordstr)
    For T = 0 To 2
        NextWord Linestr, LineLength, Wordstr, WordLength
        Param(T, L) = CDBl(Wordstr)
    Next L
    Case "Tetraloop"
'These represent especially stable tetraloops.
    'By row, sequence of the tetraloop varies.
    'FirstCol is filled with a quaternary index of the sequence.
    For L = 0 To NumTetraloops - 1
        NextLine Filestr, FileLength, Linestr, LineLength, "##"
        NextWord Linestr, LineLength, Wordstr, WordLength
        DNA_Std2Num Wordstr, NumSeq
        FirstCol(L) = 0
        For C = 0 To 3
            FirstCol(L) = FirstCol(L) * 4 + NumSeq(C)
        Next C
    Next L
    Case "DR_CalcDeltaHf(ByVal seq$)
'Function
'Calculate the association enthalpy for a given RNA sequence
'with its perfect W-C DNA complement.
'Arguments
'    Seq: The RNA sequence.
'Returns
'    Enthalpy of association, in kcal/mole.
'Notes
'    1. Standard conditions (1M Na+) is assumed.
'History

```

```

' 29-Jul-1997: From PkW's routine of the same name. PW.
Function DNA_CalcDeltaH(Byval Seq$)
    Dim Length&
    Dim B6
    Dim NumSeq()
    Dim NumSeq$(1)
    Dim Seq = LCase(Seq)

    'Create the numeric representation.
    Length = Len(Seq)
    ReDim NumSeq(0 To Length - 1)
    DNA_Str2Num Seq, NumSeq

    'length of the sequence
    'base index for traversing the sequence
    'numerical representation of the sequence

    'Lower case, please.
    'Notes
    '1. standard conditions (1M Na+) is assumed.
    '2. perfect W-C complement.

    'Sum the nearest neighbor values along the strand.
    For B = 1 To Length - 1
        DR_CalcDeltaH = DR_CalcDeltaH + DNA_DuplexH(NumSeq(B - 1), NumSeq(B))
    Next B

    'Convert to cal.
    DNA_CalcDeltaH = DNA_CalcDeltaH * 1000#
End Function

'Create the numeric representation.
Length = Len(Seq)
ReDim NumSeq(0 To Length - 1)
DNA_Str2Num Seq, NumSeq

'Initialize with initiation deltaH
DR_CalcDeltaH = DR_InitH

'Initiation with initiation deltaH
DR_CalcDeltaH = DR_CalcDeltaH + DR_InitH

'Sum the nearest neighbor values along the strand.
For B = 1 To Length - 1
    DR_CalcDeltaH = DR_CalcDeltaH + DR_DuplexH(NumSeq(B - 1), NumSeq(B))
Next B

'Convert to cal.
DR_CalcDeltaH = DR_CalcDeltaH * 1000#
End Function

'Lower case, please.
Seq = LCase(Seq)

'Create the numeric representation.
Length = Len(Seq)
ReDim NumSeq$(1)
Dim NumSeq$(0 To Length - 1)
DNA_Str2Num Seq, NumSeq$(0 To Length - 1)

'length of the sequence
'base index for traversing the sequence
'numerical representation of the sequence

'Lower case, please.
Seq = LCase(Seq)

'Create the numeric representation.
Length = Len(Seq)
ReDim NumSeq(0 To Length - 1)
DNA_Str2Num Seq, NumSeq

'Initialize with self-symmetry correction.
If Seq = DNA_RevComp(Seq) Then DNA_CalcDeltaS = DNA_CalcDeltaS + DNA_SelfS

'Add initiation term
If Instr(Seq, "c") Or Instr(Seq, "g") Then
    DNA_CalcDeltaS = DNA_CalcDeltaS + DNA_InitGcs
Else
    DNA_CalcDeltaS = DNA_CalcDeltaS + DNA_InitATs
End If

'Sum the nearest neighbor values along the strand
For B = 1 To Length - 1
    DNA_CalcDeltaS = DNA_CalcDeltaS + DNA_DuplexS(NumSeq(B - 1), NumSeq(B))
Next B

'Initialize with AT end correction.
If NumSeq(0) = 3 Then DNA_CalcDeltaH = DNA_CalcDeltaH + DNA_EndTAH

```

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```

End Function

Function DR_CalcDeltas#(ByVal seq$)
  'Calculate the association entropy for a given RNA sequence
  'with its perfect WC DNA complement.
  'Arguments
  '  seq: The sequence
  '  Entropy of association, in cal/mole/deg K.
  '  Notes
  '    1. Standard conditions (1M Na+) is assumed.
  'History
  '  - 29-Jul-1997: From PkW's routine of the same name. PW.
  'Lengths
  Dim B6          'length of the sequence
  Dim NumSeq#()   'base index for traversing the sequence
  'numerical representation of the sequence

  'Lower case, Please.
  Seq = LCase(Seq)

  'Create the numeric representation.
  Length = Len(Seq)
  RLen = NumSeq(0 To Length - 1)
  DNA_Str2Num(Seq, NumSeq)

  'Begin with initiation term.
  DR_CalcDeltas = DR_Inits

  'Sum the nearest neighbor values along the strand.
  For B = 1 To Length - 1
    DR_CalcDeltas = DR_CalcDeltas + DR_Duplexs(NumSeq(B - 1), NumSeq(B))
  Next B

  End Function

```

```

On Error Goto E           'number of sequences we are working with
Dim NumSeqs#              'index
Dim S#                     

'Determine the number of probes we are calculating TM for.
NumSeqs = UBound(Seq) + 1

'Calculate melting points
For S = 0 To NumSeqs - 1
  If S == Progress.StopAt = 0 Then Progress.CheckProgress S
  TM(S) = DNA_CalcTM(Seq(S), tmp, Conc)
Next S
Exit Sub

E: Debug.Print "Error in DNA_CalcAllTM"
Err.Raise Err.Number, , Err.Description
End Sub

Public Sub DNA_CalcClamp(Seqs(), Tclamp As CClampPars, Clamp#())
  'Function
  '  calculate the melting temperature of the Clamp of a probe
  '  to its perfect W-C complement.
  'Arguments
  '  Seq: The sequences
  '  Tclamp: An instance of the parameter class for Clamp calculations.
  'Returns
  '  Melting temperature of tightest clamp, in deg. C.
  'Notes
  '  1. Concentration is used as is, i.e. assuming the complement
  '  is present at much lower concentration.
  'On Error Goto E           'number of sequences we are working with
Dim NumSeqs#              'Indices
Dim S$, SS$                'subsequence
Dim BestTM#, ThisTM#       'current most stable clamp

'Determine the number of probes we are calculating Clamp for.
NumSeqs = UBound(Seq) + 1

'Calculate melting points.
For S = 0 To NumSeqs - 1
  If S == Progress.StopAt = 0 Then Progress.CheckProgress S
  BestTM = 0
  For SS = Tclamp.FiveP + 1 To Len(Seq(S)) - Tclamp.ThreeP
    SubSeq = Mid(Seq(S), SS, Tclamp.Length)
    ThisTM = DNA_CalcTM(SubSeq, Tclamp.Conc) + 273.15
    If ThisTM > BestTM Then BestTM = ThisTM
  Next SS
  Clamp(S) = BestTM
Next S
Exit Sub

E: Debug.Print "Error in DNA_CalcClamp"
Err.Raise Err.Number, , Err.Description
End Sub

```

```

Public Sub DR_CalcClamp (Seqs(), TClamp As CClampPars, Clamp#())
'-----'
'Function
'Calculate the melting temperature of the Clamp of a probe
'    to its perfect RNA W-C complement.
'Arguments
'    Seq: The sequences
'    TClamp: An instance of the parameter class for Clamp calculations.
'Returns
'Notes
'    Melting temperature of tightest clamp, in dgC.
'    1. Concentration is used as is, i.e. assuming the complement
'    is present at much lower concentration.
'-----'
On Error GoTo E
Dim NumSeqs      'number of sequences we are working with
Dim S$, SS$      'indices
Dim SubSeqs$     'subsequence
Dim BestTM#, ThisTM#   current most stable clamp
'Determine the number of probes we are calculating Clamp for.
NumSeqs = UBound(Seq) + 1
'Calculate melting points.
For S = 0 To NumSeqs - 1
    If S = Progress.StopAt = 0 Then Progress.CheckProgress S
    BestTM = 0
    For SS = TClamp.FiveP + 1 To Len(Seq(S)) - TClamp.ThreeP - TClamp.Length + 1
        SubSeq = Mid(Seq, SS, TClamp.Length)
        ThisTM = DR_CalcTM(SubSeq, TClamp, Conc) + 273.15
        If ThisTM > BestTM Then BestTM = ThisTM
    Next SS
    Clamp(S) = BestTM
Next S
Exit Sub
E: Debug.Print "Error in DR_CalcClamp"
Err.Raise Err.Number, , Err.Description
End Sub
Public sub DNA_CalcDGH (Seqs(), dGHP As CDGHPars, dGH#())
'-----'
'Function
'Calculate the hairpin dGs of an array of oligos.
'Arguments
'    Seq: The sequences.
'    dGHP: An instance of the parameter class for dGH calculations.
'    dGH: The hairpin dGs.
'-----'
On Error GoTo E
Dim NumSeq#()    'numeric representation of the sequence
Dim S#             'index
'Recalculate all G parameter matrices at current temperature.

```

```

CalcGFromH 4, DNahp_StackH, DNahp_StackS, DGHP_T, DNahp_StackG
CalcGFromH 4, DNahp_TstackH, DNahp_TstackS, DGHP_T, DNahp_TstackG
CalcGFromH 4, DNahp_DangleH, DNahp_Dangles, DGHP_T, DNahp_DangleG
CalcGFromH 2, DNahp_Loops, DGHP_T, DNahp_Loops, DGHP_T, DNahp_LoopG
CalcGFromH 1, DNahp_TloopH, DNahp_TloopS, DGHP_T, DNahp_TLoopG
'Calculate dGs.
For S = 0 To UBound(seq)
    If S = Progress.StopAt = 0 Then Progress.CheckProgress S
    ReDim NumSeq(0 To Len(Seq(S)) - 1)
    DNA_ST2Num Seq(S), NumSeq
    dG(S) = DNA_BestHairpin(NumSeq)
Next S
Exit Sub
E: Debug.Print "Error in DNA_CalcGH"
Err.Raise Err.Number, , Err.Description
End Sub
Public Sub DNA_CalcDGP (Seq$, dGDP As CDGDPars, dGD#())
'-----'
'Function
'Calculate the duplex dGs of an array of oligos.
'Arguments
'    Seq: The sequences.
'    dGDP: An instance of the parameter class for dGD calculations.
'    dGD: The hairpin dGs.
'-----'
On Error GoTo E
Dim NumSeq#()    'numeric representation of the sequence
Dim S#             'index
'Calculate dGs.
For S = 0 To UBound(Seq)
    If S = Progress.StopAt = 0 Then Progress.CheckProgress S
    dG(S) = (DNA_CalcDeltaS(Seq(S))) / 1000#
    DNA_CalcDeltaS(Seq(S)) = (dGDP.T + 273.15) *
    DNA_CalcDeltas(Seq(S))
Next S
Exit Sub
E: Debug.Print "Error in DNA_CalcDGP"
Err.Raise Err.Number, , Err.Description
End Sub
Public Sub DR_CalcDD (Seqs(), dGDP As CDGDPars, dGD#())
'-----'
'Function
'Calculate the duplex dGs of an array of oligos.
'Arguments
'    Seq: The sequences.
'    dGDP: An instance of the parameter class for dGD calculations.
'    dGD: The hairpin dGs.
'-----'
On Error GoTo E
Dim NumSeq#()    'numeric representation of the sequence
Dim S#             'index
'Calculate dGs.

```

```

For S = 0 To UBound(Seq)
  If S - Progress.StopAt = 0 Then Progress.CheckProgress S
  / 1000#
  dGf(S) = fDR_CalcDeltaH(Seq(S)) - (dGDP.T + 273.15) * DR_CalcDeltas(Seq(S))
Next S
Exit Sub
E: Debug.Print "Error in DNA CalcGDF"
Err.Raise Err.Number, , Err.Description
End Sub

Public Sub DNA_CalcGDM(Seqs(), dGMP As cdGMPars, dGM#())
  Dim S
  'Function
  'Calculate the MFold dGs of an array of oligos.
  'Arguments
  '  Seq: The sequences.
  '  dGMP: An instance of the parameter class for dGM calculations.
  '  dGM: The hairpin dGs.
  '  On Error Goto E
  Dim S6   'Index
  'Calculate dGs.
  For S = 0 To UBound(Seq)
    If S - Progress.StopAt = 0 Then Progress.CheckProgress S
    dGM(S) = frmMain.MFold.Seq(S), Val(dGMP.T), True
  Next S
  Exit Sub
  E: Debug.Print "Error in DNA_CalcGDM"
  Err.Raise Err.Number, , Err.Description
End Sub

Public Sub RNA_CalcGDM(Seqs(), dGHP As cdGHPars, dGH#())
  Dim S
  'Function
  'Calculate the hairpin dGs of an array of oligos.
  'Arguments
  '  Seq: The sequences.
  '  dGHP: An instance of the parameter class for dGH calculations.
  '  dGH: The hairpin dGs.
  '  On Error Goto E
  Dim NumSeqs() 'numeric representation of the sequence
  Dim S6   'Index
  'Recalculate all G parameter matrices at current temperature.
  CalcGFromHS 4, RNahp_StackH, RNahp_StackS, dGHP.T, RNahp_stackG
  CalcGFromHS 4, RNahp_TStackH, RNahp_TStackS, dGHP.T, RNahp_TstackG
  CalcGFromHS 4, RNahp_DangleH, RNahp_Dangles, dGHP.T, RNahp_DangleG
  CalcGFromHS 2, RNahp_LoopH, RNahp_Loops, dGHP.T, RNahp_LoopG
  CalcGFromHS 1, RNahp_TLoopH, RNahp_TLoops, dGHP.T, RNahp_TLoopG
  'Calculate dGs.

```

```

For S = 0 To UBound(Seq)
  If S - Progress.StopAt = 0 Then Progress.CheckProgress S
  Redim NumSeq(0 To Len(Seq(S)) - 1)
  DNA_ST(2)Num Seq(S), NumSeq
  dGH(S) = RNA_BestHairpin(NumSeq)
Next S
Exit Sub
E: Debug.Print "Error in RNA_CalcGHD"
Err.Raise Err.Number, , Err.Description
End Sub

Public Sub RNA_CalcGHD(Seqs(), dGM#())
  Dim S
  'Function
  'Calculate the MFold dGs of an array of oligos.
  'Arguments
  '  Seq: The sequences.
  '  dGM: An instance of the parameter class for dGM calculations.
  '  dGM: The MFold dGs.
  '  On Error Goto E
  Dim S6   'Index
  'Calculate dGs.
  For S = 0 To UBound(Seq)
    If S - Progress.StopAt = 0 Then Progress.CheckProgress S
    dGM(S) = frmMain.MFold.Seq(S), Val(dGMP.T), False
  Next S
  Exit Sub
  E: Debug.Print "Error in RNA_CalcGHD"
  Err.Raise Err.Number, , Err.Description
End Sub

Public Sub DR_CalcAllTM(Seqs(), tmp As cdMPars, TM#())
  Dim S
  'Function
  'Calculate the melting temperature of an array of DNA probes with their
  'perfect W-C RNA target complements.
  'Arguments
  '  Seq: The sequences.
  '  TM: An instance of the parameter class for TM calculations.
  '  Returns
  '    Melting temperature, in deg. C.
  '  Notes
  '    1. Concentration is used as is, i.e. assuming the complement
  '       is present at much lower concentration.
  'History
  '  29-Jul-1997: From PKW's routine of the same name. PW.
  On Error Goto E
  Dim NumSeqs
  Dim S6   'Index
  'number of sequences we are working with
  'Determine the number of probes we are calculating TM for.
  NumSeqs = UBound(Seq) - LBound(Seq) + 1

```

```

'Calculate melting points
For S = 0 To NumSeqs - 1
    If S = Progress.StopAt = 0 Then Progress.CheckProgress S
    TM(S) = DR_CalcTM(Seq(S), tmp.conc)
Next S
Exit Sub
B: Debug.Print "Error in DR_CalcAllTM"
Err.Raise Err.Number, , Err.Description
End Sub

Public Function DR_CalcTM#(Seqs$, conc$)
    'Function
    ' Calculate the TM of one DNA/RNA duplex. Sequence given is DNA.
    ' Calculate the TM of one DNA/DNA duplex.
    DR_CalcTM = DR_CalcDeltaH(seq) / (DR_CalcDeltas(seq) + RGas * Log(conc)) -
    273.15
End Function

Public Function DNA_CalcTM#(seq$, conc$)
    'Function
    ' Calculate the TM of one DNA/DNA duplex.
    DNA_CalcTM = DNA_CalcDeltaH(seq) / (DNA_CalcDeltas(seq) + RGas * Log(conc)) -
    273.15
End Function

Attribute VB_Name = "Utilities"
Option Explicit.

'A quicksort routine for doubles, based on NR code.
Private Declare Function vbSort2 Lib "vb5ndll.dll" -
    (ByVal N%, ByRef Vector!, ByRef Index%) As Long

Public sub Calcrun(Pos%(), RP As crunPars, Run%())
    'Function
    ' Calculate the runs in a set of positions.
    'Arguments
    'Pos: The positions.
    'RP: The run parameters.
    'Run: The returned run.
    'Method
    'Begin by looping over all positions, extending down over all consecutive
    'entries, then up over all consecutive entries. Then mark in the active
    'position the length of run found. Next, eliminate all runs that are
    'too short. Finally, for each run, mark the requested number of members
    'of the run, at the requested spacing, as being members. If the run is
    'shorter than can accommodate the requested number of members, fill in as
    'many as possible, while still preserving the requested spacing.
    'Notes
    '1. Run must be allocated and sized correctly by the caller.
    'Index
    Dim Pos%
    Dim EndUp%
    Dim EndDown%

```

```

    Dim StepDirk      'current step direction
    Dim M%           'index
    'Loop over positions.
    For P = 0 To UBound(Pos)
        EndDown = P
        Do While ((P - EndDown) = (Pos(P) - Pos(EndDown)))
            EndDown = EndDown - 1
            If EndDown = -1 Then Exit Do
        Loop
        EndUp = EndDown + 1
        EndUp = P
        Do While ((EndUp - P) = (Pos(EndUp) - Pos(P)))
            EndUp = EndUp + 1
            If EndUp = UBound(Pos) + 1 Then Exit Do
        Loop
        EndUp = EndUp - 1
        EndUp = EndUp - EndDown + 1
        Run(P) = EndUp - EndDown + 1
    Next P

    'Remove runs that are too short.
    For P = 0 To UBound(Run)
        If Run(P) < RP.Min Then Run(P) = 0
    Next P

    'Pick out requested elements from each run.
    P = 0
    Do While P < UBound(Run)

        'Find next run.
        Do While Run(P) = 0 And P < UBound(Run)
            P = P + 1
        Loop

        'Record the ends, step to the middle (to the right of middle for even
        'lengths).
        EndDown = P
        Do While Run(P) = 0 And P < UBound(Run)
            P = P + 1
        Loop

        'Mark elements, stepping down from the middle first. This ensures that all
        'elements get marked for even lengths, spacing=1.
        StepDirk = -1
        For M = 1 To RP.Num
            Run(P) = -Run(P)
            P = Fix((EndDown + EndUp + 1) / 2)
            P = P + StepDirk * M * RP.Spacing
            If (P < EndDown Or P > EndUp) Then Exit For
            StepDirk = StepDirk * -1
        Next M

        'Move over this run.
        P = EndUp + 1
    Loop

    'Convert marked elements back to run length.
    For P = 0 To UBound(Run)
        If Run(P) > 0 Then Run(P) = 0
        If Run(P) < 0 Then Run(P) = -Run(P)
    Next P

```

```

Next P
End Sub

Public Function IsGoodRS(RS As Recordset) As Boolean
    'Function
    ' Check whether it is possible to access the records of
    ' recordset connected to the sequence, probaset, or probeset table.
    ' No current record need exist for this function to return true.
    '-----'
    On Error GoTo Err
    IsGoodRS = False
    If IsNull(RS) Then Exit Function
    If (RS.EOF = True And RS.BOF = True) Then Exit Function
    IsGoodRS = True
    Err:
    End Function

    Public Function NumRecords(RS As Recordset)
        '-----'
        ' Function
        ' Count the number of records in the recordset.
        '-----'
        ' Notes
        ' A side effect of this routine is to position the recordset at the
        ' first record.
        '-----'
        RS.MoveLast
        RS.MoveFirst
        NumRecords = RS.RecordCount
    End Function

    Public Sub UnPackSequence(ByRef Seqstr As String, ByRef NumStr As String)
        'Seqstr is assumed to hold a packed sequence.
        Dim NewSeqStr$,
        Dim Bases, I&
        NumStr = ""
        Bases = 1
        Do While Len(Seqstr) - Bases > 60
            NumStr = NumStr & Format(Bases, "#####"), "#####") & vbCrLf
            For I = Bases To Bases + 59
                NewSeqstr = NewSeqstr & Mid(Seqstr, I, 1)
                If (I Mod 10 = 0) Then NewSeqstr = NewSeqstr & " "
            Next I
            Bases = Bases + 60
            NewSeqstr = NewSeqstr + vbCrLf
        Loop
        If Bases <> Len(Seqstr) Then
            NumStr = NumStr & Format(Bases, "#####"), "#####")
            For I = Bases To Len(Seqstr)
                NewSeqstr = NewSeqstr & Mid(Seqstr, I, 1)
                If (I Mod 10 = 0) Then NewSeqstr = NewSeqstr & " "
            Next I
            If NewSeqstr = Seqstr Then
                End If
                Seqstr = NewSeqstr
            End Sub
    End If

```

```

Public Function StrFields(str$, lineLen$, ByVal Fieldlen$, Optional Extra$)
    'Returns leading field f a string, and shortens the string
    'by the fieldlength + extra.
    'If the line is too short, as much of the requested field as possible
    'will be returned.

    If Fieldlen > lineLen Then Fieldlen = lineLen
    StrField = Trim(Left$(str$, Fieldlen))
    lineLen = lineLen - Fieldlen
    If Not IsMissing(Extra) Then
        LineLen = lineLen - Extra
        If lineLen < 0 Then lineLen = 0
    End If
    Str = Right$(str$, lineLen)
    End Function

    Public Sub NextWord(str$, StrLen$, Words$, WordLen$, Optional Extra$)
        'strips leading spaces, copies leading word, reduces lineLength
        Do While ((Mid$(str$, 1, 1) = " ") And (Strlen > 0))
            Strlen = Strlen - 1
            Str = Right$(str$, Strlen)
        Loop
        If Instr$(str$, " ") <> 0 Then
            WordLen = Instr$(str$, " ")
            Strlen = Strlen - 1
        Else
            WordLen = Strlen
        End If
        If WordLen <> 0 Then
            Word = Mid$(str$, 1, WordLen)
            Strlen = Strlen - WordLen
            Str = Right$(str$, Strlen)
        Else
            Word = ""
            Str = ""
            Strlen = 0
        End If
    End Sub

    Public Sub NextLine(Filestr$, FileLen$, Linestr$, lineLen$, CommentChar$)
        'Return the next line that doesn't begin with CommentChar
        lineLen = 0
        Do While (lineLen = 0 And FileLen > 0)
            lineLen = Instr(Filestr, vbCrLf) - 1
            Linestr = Left$(Filestr, lineLen)
            Filelen = Filelen - lineLen - 2
            Filestr = Right$(Filestr, Filelen)
            If (Mid$(Linestr, 1, 1) = CommentChar) Then lineLen = 0
        Loop
    End Sub

    Public sub Quicksort(x(), Index(), LB$, UB$)
        'Quicksorts the array X into a ascending order,
        'simultaneously sorting I to provide a sort index.
        Dim N$, foo
        N = UB - LB + 1
        foo = vb5Sort2(N, X(LB), Indax(LB))

```

```

End Sub

Public Function BinarySearch$(Vector(), Value$, Ut, Lt)
'Function
' Find the index of a value in a sorted array.
Do While Ut <= Lt
    BinarySearch = (Ut + Lt) / 2
    If Vector(BinarySearch) = Value Then Exit Function
    If Vector(BinarySearch) > Value Then
        Ut = BinarySearch
    Else
        Lt = BinarySearch
    End If
Loop
End Function

Public Function PackSequence$(SeqStr$)
Dim NewStr$, Ch$,
    For I = 1 To Len(SeqStr)
        Ch = Mid$(SeqStr, I, 1)
        If Instr("ACGTUCGt", Ch) Then PackSequence = PackSequence & Ch
    Next I
End Function

```